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(54) Title: A METHOD FOR TREATING HERPES VIRUSES

(57) Abstract: The present invention relates to a method for selecting an anti-herpes viral compound and a method for selectively inhibiting herpesvrus in a human host in need of such treatment. The present invention relates to a method for selecting an anti-herpes viral compound and a method for selectively inhibiting herpsvrus in a human host in need of such treatment.

A METHOD FOR TREATING HERPES VIRUSES

FIELD OF THE INVENTION

The present invention relates to a method for selecting an anti-herpes viral compound and a method for selectively inhibiting herpes viruses in a human host in need of such treatment.

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BACKGROUND OF THE INVENTION

The herpesviruses comprise a large family of double stranded DNA viruses. Eight of the herpes viruses, herpes simplex virus types 1 and 2 (HSV-1 and HSV-2), varicella zoster virus (VZV), human cytomegalovirus (HCMV), Epstein-Barr virus (EBV), and human herpes viruses 6, 7, and 8 (HHV-6, HHV-7, and HHV-8), have been shown to infect humans. Several of these viruses are important human pathogens.

HSV-1 is estimated to affect 100 million people in the U.S. Primary infection of HSV-1 usually occurs between the ages of one and four. Cold sores, the visible symptom, typically appear at a later age, with 20-45% of the population over the age of fifteen affected (Whitley, Clin. Intect. Dis., 26:541-555, 1998).

Genital herpes (HSV-2) is the second most common sexually transmitted disease, with approximately 22% of the U.S population infected with this virus (Fleming 1997).

VZV is the causative agent of chicken pox upon primary infection and can recur in adults as zoster.

EBV results in approximately two million cases of infectious mononucleosis in the U.S. each year. It can also cause lymphomas in immunocompromised patients and has been associated with Burkitt's lymphoma, nasopharyngeal carcinoma, and Hodgkins disease.

Infection with HCMV often occurs during childhood and is typically asymptomatic except in immunocompriomised patients where it causes significant morbidity and mortality.

HHV-6 is the causitive agent of roseola and may be associated with multiple sclerosis and chronic fatigue syndrome. HHV-7 disease association is unclear, but it may be involved in some cases of roseola. HHV-8 has been associated with Karposi's sarcoma, body cavity based lymphomas, and multiple myeloma.

These viruses are capable of residing in a latent state within the host. Reactivation of latent virus results from response to environmental stimuli (ex. UV exposure, stress,

etc.). Infections or recurrence can be life threatening in immunocompromised patients such as AIDS or transplant patients where HCMV can result in retinitis, pneumonia, and gastrointestinal disease.

The increased immunocompromised population has created an unmet medical need for antivirals against herpesviruses because current therapies do not have a sufficiently broad spectrum against this family of viruses and/or they have limited utility due to toxicity. The present invention provides a method for selectively inhibiting herpesviruses DNA polymerase with compounds that have broad spectrum activity. The method offers a distinct advantage in the treatment of patients in need, particularly immunocompromised patients at risk of infection or reactivation by many members of the herpesvirus family.

SUMMARY OF THE INVENTION

The present invention provides a method of selecting compounds that inhibit herpes viruses comprising:

- 15 a) measuring IC₅₀ of a compound of interest that inhibits a wild type herpes virus,
 - b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant herpes virus which is the same strain of the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and

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- d) selecting the compound of interest wherein the IC₅₀ of step b is at least 3 times greater than the IC₅₀ of step a.
 - In above method, the order of step a and step b are interchangeable.

The present invention further provides a method of selecting compounds that inhibit herpes viruses comprising:

- a) measuring IC₅₀ of a compound of interest that inhibits a wild type HSV-1,
- 25 b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant HSV-1 which is the same strain of the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - d) selecting the compound of interest wherein the IC₅₀ of step b is at least 3 times greater than the IC₅₀ of step a.
- In above method, the order of step a and step b are interchangeable.

The present invention further provides a method of selecting compounds that inhibit herpes viruses comprising:

a) measuring IC₅₀ of a compound of interest that inhibits a wild type HSV-2,

b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant HSV-2 which is the same strain of the wild type herpes virus,

c) comparing IC₅₀ of step a with IC₅₀ of step b; and

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d) selecting the compound of interest wherein the IC_{50} of step b is at least 3 times greater than the IC_{50} of step a.

In above method, the order of step a and step b are interchangeable.

The present invention further provides a method of selecting compounds that inhibit herpes viruses comprising:

- a) measuring IC₅₀ of a compound of interest that inhibits a wild type HCMV,
- measuring IC₅₀ of the same compound that inhibits a binding domain mutant HCMV which is the same strain of the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - d) selecting the compound of interest wherein the IC_{50} of step b is at least 3 times greater than the IC_{50} of step a.

In above method, the order of step a and step b are interchangeable.

The present invention further provides a method for selectively treating diseases caused by herpes viruses in a human host comprising administering a compound to a human in need of such treatment wherein said compound inhibits herpes viruses by interaction with the binding domain in the viral DNA polymerase.

The present invention further provides method for selectively inhibiting herpes viruses in a human host comprising administering a compound to a human in need of such treatment wherein IC₅₀ of the compound that inhibits a binding domain mutant herpes virus is at lease 3 times greater than IC₅₀ of the compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes virus.

The present invention further provides a compound for treating herpesviral infections in a human host wherein IC₅₀ of the compound that inhibits a binding domain mutant herpes virus is at lease 5 times greater than IC₅₀ of the compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes virus.

The present invention further provides a compound for treating herpesviral infections in a human host wherein said compound inhibits the herpesvirus by interacting with the binding domain in the viral DNA polymerase.

The present invention further provides a compound for the inhibiting of herpesvirus DNA polymerases wherein serial passage of a wild type herpes virus in the presence of said

compound results in a change of the wild type HSV-1 polymerase at amino acid 823 from valine to alanine.

The present invention further provides a compound for inhibiting herpesvirus DNA polymerases wherein serial passage of a wild type herpes virus in the presence of said compound results a change of the wild type HCMV polymerase at amino acid 823 from valine to alanine and at amino acid 824 from valine to leucine.

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The present invention further provides a mutant herpesvirus DNA molecule having a nucleotide sequence selected from a group consisting of SEQ.ID.NO. 1; SEQ.ID.NO. 3; SEQ.ID.NO. 5; SEQ.ID.NO. 7; SEQ.ID.NO. 9; and SEQ.ID.NO. 11.

The present invention further provides a mutant herpesvirus polymerase amino acid molecule having an amino acid sequence selected from a group consisting of SEQ.ID.NO. 2; SEQ.ID.NO. 4; SEQ.ID.NO. 6; SEQ.ID.NO. 8; SEQ.ID.NO. 10 and SEQ.ID.NO. 12.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 – examples of 4-oxo-DHQ and 4-oxo-DHTP compounds.

Figure 2 - Herpesvirus' polymerases amino acid conserved region.

Figure 3 – Recovered virus after serial passage of HSV-1 in presence of 20 μM of compound No. 17.

Figure 4 – Comparision of Wild HSV-1 and HSV-2 herpesvirus DNA polymerase amino acid sequences alligned by amino acid homology. (Seq. No: 14-19)

Figure 5 - Mutant Herpes Virus DNA and amino acid sequence list. (Seq. No: 1-12)

Figure 6 – Wild HCMV herpesvirus DNA polymerases amino acid sequence. (Seq. No 13)

DETAILED DESCRIPTION OF THE INVENTION

A key enzyme in the replication of all herpesviruses is the virus-coded DNA polymerase. Most of the currently available anti-herpes drugs target the viral DNA polymerase. Drugs such as Foscarnet acts by direct inhibition of the viral polymerase. These drugs are non-nucleoside inhibitors of herpesvirus DNA polymerases. Others such as the nucleoside analogs, Acyclovir, Penciclovir and Ganciclovir must first be phosphorylated to the monophosphate forms by virus encoded kinases and, further phosphorylated to triphosphate by cellular enzymes before they are active inhibitors. The triphosphate forms of these nucleoside analogs inhibit polymerases by competing with the binding of natural

triphosphates and their subsequent insertion into growing DNA strands. These drugs are known as nucleoside inhibitors of herpesvirus DNA polymerases.

One of the limitations of the currently available drugs is that they are active against only a few of the eight human herpesviruses. For example, Acyclovir and Penciclovir inhibit HSV and VZV replication but have poor activity against CMV.

In order to identify antiviral compounds that would have the potential to inhibit replication of most of the human herpesviruses, compounds are *in vitro* screened for inhibitors of herpesvirus DNA polymerase activity. Because portions of the amino acid sequence of the polymerases are highly conserved within the herpesvirus family it is possible to discover small molecules that inhibit herpesvirus polymerases but not cellular DNA polymerases. Using this biochemical approach, several new classes of compounds such as the 4-hydroxyquinoline derivatives (4-HQ), 4-oxo-dihydroquinoline derivatives (4-oxo-DHQ) and 4-oxo-dihydrothienopyridine derivatives (4-oxo-DHTP) were discovered as potent, non-nucleoside herpesvirus DNA polymerase inhibitors. *In vitro* polymerase assays and/or *in vivo* cell culture assays have demonstrated that these compounds inhibit HSV-1, HSV-2, HCMV, VZV, EBV, and HHV-8 replication.

4-Oxo-DHQ and 4-oxo-DHTP are derivatives of formula I

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wherein ring A is a saturated or unsaturated fused double or triple heterocyclic ring having 1, 2, 3 or 4 heteroatoms selected from group consisting of oxygen, sulfur, or nitrogen; and wherein R and X are the appropriated substitutents, respectively.

Examples of 4-HQ compounds, 4-oxo-DHQ compounds and 4-oxo-DHTP compounds are illustrated in **Figure 1**.

Antiviral activity of these examples are shown in Table 1 below. As shown in Table 1, these compounds inhibit HSV-1 and HSV-2 as well or better than the current commercially available drug Acyclovir.

Table 1
Antiviral Activity of 4-oxo DHQ/4-oxo DTHP Against HSV-1 and HSV-2

	Compound IC ₅₀ (uM)								
virus	1	2	3	4	5	ACV			
HSV-1 KOS	2.0	3.8	3.2	3.2	3.3	3.6			
HSV-1 F	2.5	2.3	2.2	2.1	2.6	1.3			
HSV-1 DJL	2.5	2.6	1.8	2.2	2.7	1.8			
HSV-1 Patton	ND	5.3	7.7	4.3	10	9.3			
HSV-2 MS	2.0	2.5	2.8	2.5	2.5	10			
HSV-2 35D	ND	5.4	5.0	3.2	8.1	6.0			
HSV-2 186	2.0	2.3	3.2	2.3	4.2	>10			

It has also been discovered that point mutations within the HSV-1 polymerase gene that confer resistance to Acyclovir and other nucleoside analogs do not result in resistance to the 4-HQ, 4-oxo-DHQs or 4-oxo-DHTPs. Serial passage of wild type HSV-1 in the presence of 4-oxo-DHQ results in the isolation of mutants that are highly resistant (>20 fold increase in the IC₅₀) to these compounds while retaining sensitivity to nucleoside inhibitors such as Acyclovir.

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In order to determine the mechanism of action of 4-HQ, 4-oxo-DHQ and 4-oxo-DHTP compounds against herpes viruses, mutants resistant to these compounds are isolated by serial passage of the virus in the presence of a 4-oxo-DHQ compound. Sequencing analysis of HSV-1 and HSV-2 strains resistant to the 4-oxo-DHQ identifies that HSV-1 (KOS strain) polymerase protein and its homologous HSV-2 have a conserved region (a binding domain), which is a critical contact point for these compounds. While amino acid numbering of the DNA polymerase may vary between strains of HSV-1 and HSV-2, this binding domain encompassing the HSV-1 (KOS) strain amino acid 823 is highly conserved in herpesviruses and can be identified by alligning the homologous amino acids of this domain as shown in Fig 2.

In HSV-1 and HSV-2 strains resistant to the 4-oxo-DHQ and similar compounds, a change of valine to an alanine at the binding domain provides full resistance.

In the HSV-1 DNA polymerase, resistance is also found when a valine changes to methionine at amino acid 823 but only when accompanied by a second amino acid change.

Isolation of HCMV resistant to 4-oxo-DHQ's is found to be very difficult. Comparison of the amino acid sequence of the HSV polymerase (Y-G-F-T-G-V-Q-H-G) and HCMV polymerase (Y-G-F-T-G-V-V-N-G) in the region of amino acid 823 (underlined amino acid) shows that there is a second value at position 824 in the HCMV

polymerase. In vitro assay using mutant HCMV polymerases demonstrates that full resistance to the 4-oxo-DHQs requires changes at both amino acids 823 (a valine to alanine) and 824 (a valine to leucine). A HCMV polymerase gene containing V823A and V824L mutations is used in marker rescue experiments to generate a viral mutant. This mutant has an IC₅₀ approximately 7-fold above that of wild-type HCMV.

The HSV-1, HSV-2 and HCMV mutants are also found to be resistant to other non-nucleoside inhibitors such as the 4-oxo-DHTP and similar compounds. However, when the binding domain mutants (e. g. HSV-1 V823A, HSV-2-MS V826A, HSV-2-186 V828A, and HCMV V823A/V824L mutants) are tested in plaque reduction assays against a series of nucleoside polymerase inhibitors and the non-nucleoside inhibitor such as Foscarnet, replication of the mutants is found to be inhibited by all of the currently marketed anti-herpes polymerase inhibitors tested.

These studies demonstrate that certain non-nucleosides like 4-HQ, 4-oxo-DHQ and 4-oxo-DHTP compounds bind to a different site on the herpes polymerase than the nucleoside inhibitors and Foscarnet. The valine at the binding domain is conserved in the DNA polymerases of six of the eight human herpesviruses and several animal herpesviruses, and appears to play a critical role in the antiviral activity of the 4-HQ, 4-oxo-DHQ and 4-oxo-DHTP compounds. (See Figure 2)

Since mutation at the binding domain negates these non-nucleoside inhibitors' activities, compounds could be tested against wild type polymerases and the mutant polymerases to establish the probability of similar binding. We refer to this property of compounds as interaction with the binding domain. Since compounds that interact with the binding domain have exhibited broad-spectrum activity against herpesviruses, this invention provides a method for selecting compounds to treat individuals such as immunocompromised patients who are afflicted with multple herpesvirus infections.

Definitions

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The term "wild-type" refers to a gene or gene product which has the characteristics of that gene or gene product when isolated from a naturally occurring source. A wild-type gene is that which is most frequently observed in a population and is thus arbitrarily designated the "normal" or "wild-type" form of the gene.

In contrast, the term "mutant" refers to a gene or gene product which displays modifications in sequence and or functional properties (i.e., altered characteristics) when

compared to the wild-type gene or gene product. It is noted that naturally-occurring mutants can be isolated; these are identified by the fact that they have altered characteristics when compared to the wild-type gene or gene product.

IC₅₀ refers to concentration of a drug that inhibits virus growth by 50%.

Wild type HSV-1 and HSV-2 strains are listed in Figure 4.

Wild type HCMV is listed in SEQ. ID. NO.13.

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The term "Iudr" refers to antiviral drug Iododeoxyuridine.

The term "Bvdu" refers to antiviral drug Bromovinyldeoxyuridine.

The term "ACV" refers to antiviral drug Acyclovir.

The term "AraC" refers to antiviral drug Arabinosylcytidine.

The term "AraT" refers to antiviral drug Arabinosylthymine.

The term "AraA" refers to antiviral drug Arabinosyladenine.

The term "GCV" refers to antiviral drug Ganciclovir.

The term "CDV" refers to antiviral drug Cidofovir.

The term "PFA" refers to antiviral drug Foscarnet.

The term "binding domain" refers to a conserved region in herpesvirus DNA polymerases. The herpesvirus DNA polymerases have seven (7) conserved regions. The binding domain is within the thrid conserved region (see Figure 2). When the binding domain contacts with an inhibitor, at least one amino acid in the binding domain mutates and provides the resistance. In general, the binding domain is at an amino acid sequence position 818-829 of the HSV-1 DNA polymerase or the homologous region in other herpes virus DNA polymerases (see Figure 2).

The term "a binding domain mutant herpes virus" refers to a herpes virus containing a binding domain mutation.

More specifically, the binding domain in HSV-1 strains, KOS, F, DJL and Patton are at amino acid sequence position 823. The binding domain in HSV-2 MS-M1 strain is at amino acid sequence position 826. The binding domain in HSV-2 186 strain is at amino acid sequence position 828. The binding domain in HCMV AD 169 strains is at amino acid sequence position 823-824.

The term "XxxxY" refers to an amino acid sequence position xxx, a single amino acid X in wild type is changed to an amino acid Y.

For example, the term "V823A" refers to an amino acid sequence position 823, a Valine found in wild type is changed to alanine in mutant strain.

The term "V824L" refers to an amino acid sequence position 824, a Valine found in wild type is changed to Leucine in mutant strain.

The term "V826A" refers to an amino acid sequence position 826, a Valine found in wild type is change to alanine in mutant strain.

The term "V828A" refers to an amino acid sequence position 828, a Valine found in wild type is change to alanine in mutant strain.

A table of amino acids and their representative abbreviations, symbols and codons is set forth below in the following Table.

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Amino acid	Abbrev.	Symbol	<u> </u>			don(s)		
Alanine	Ala	A	GCA	GCC	GCG	GCU		
Cysteine	Cys	C	UGC	UGU				
Aspartic acid	Asp	D	GAC	GAU				
Glutamic acid	Glu	E	GAA	GAG				
Phenylalanine	Phe	F	UUC	טטט				
Glycine	Gly	G	GGA	GGC	GGG	GGU		
Histidine	His	H	CAC	CAU				
Isoleucine	Ile	I	AUA	AUC	AUU			
Lysine	Lys	K	AAA	AAG				
Leucine	Leu	L	UUA	UUG	CUA	CUC	CUG	CUU
Methionine	Met	M	AUG					
Asparagine	Asn	N	AAC	AAU				
Proline	Pro	P	CCA	CCC	CCG	CCU		
Glutamine	Gln	Q	CAA	CAG				
Arginine	Arg	R	AGA	AGG	CGA	CGC	CGG	CGU
Serine	Ser	S	AGC	AGU	UCA	UCC	UCG	UCU
Threonine	Thr	T	ACA	ACC	ACG	ACU		
Valine	Val	V	GUA	GUC	GUG	GUU		
Tryptophan	Trp	W	UGG					
Tyrosine	Tyr	Y	UAC	UAU				

MATERIALS AND METHODS

Cell and Viruses

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African green monkey kidney cells (Vero) and human foreskin fibroblast cells (HFF) and herpes viruses can be obtained from the American Type Culture Collection (ATCC). Media is defined as Dulbecco's modified Eagle media (DMEM) containing 10% fetal bovine serum (FBS) and supplemented with antibiotics. Cells are maintained in media at 37°C in a humidified atmosphere of 5% CO². HSV-1 strains F, Patton and DJL, HSV-2 strains MS, 35D and 186, and HCMV strain AD169 are used in these studies. Strain DJL is a clinical isolate of HSV-1 isolated in our lab from a primary oral lesion.

Measuring IC₅₀ of a Compound of Interest That Inhibits Herpes Viruses

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Preparation of Virus Stocks: HSV-1 and HSV-2 stocks are grown in Vero cells. HCMV stocks are grown in HFF cells. Approximately 1 ml of media containing sufficient virus to infect approximately 0.1% to 1% of the cells (multiplicity of infection of 0.001 to 0.01 PFU/cell) is added to a T-150 cell culture flask containing a confluent monolayer of cells. The cells are incubated at 37°C for approximately 1 hour. Approximately 50 ml of media is then added to the flask and the cells are incubated at 37°C until viral cytopathic effect (cpe) is apparent in 100% of the cells. The flask is then placed at -80°C for at least 30 min. The flask containing frozen media and cells is placed in a 37°C water bath until the media is thawed. This process disrupts the cells and releases virus into the media. 1 ml aliquots of media containing virus are dispensed into tubes and stored at -80°C. These aliquots of media containing virus are referred to as virus stocks.

Titrating Virus Stocks: Aliquots of virus are thawed at 37°C and serially diluted (10 fold dilutions) in media. 0.1 ml of each dilution of virus is placed in a single well of 24-well cell culture dish containing a confluent monolayer of cells (Vero cells for HSV-1 and HSV-2, HFF cells for HCMV) and incubated at 37°C for 1 h. The virus innoculum is then removed and 1 ml of media containing 0.8% carboxymethylcellulose (CMC) is added to each well of the dish. The dish is incubated at 37°C for approximately 2-3 days (HSV-1 and HSV-2) or 6-9 days (HCMV) to allow sufficient growth of virus to form plaques in the cell monolayer. Plaques can be observed and counted microscopically or by staining the cells with 0.1% crystal violet in 20% ethanol. The virus titer which is expressed as plaque forming units (PFU) per ml is obtained by counting the plaques in a well and correcting for the dilution of the viral innoculum.

Plaque Reduction Assays: Antiviral activity of compounds against herpesviruses such as HSV-1, HSV-2, or HCMV can be measured using plaque reduction assays. 0.1 ml of media containing approximately 50 PFU of virus is added to each well of a 24-well cell culture dish containing a confluent monolayer of cells (Vero cells for HSV-1 and HSV-2, HFF cells for HCMV). Compounds are dissolved in 100% DMSO and diluted in 100% DMSO as 200x stocks of the desired final drug concentration. Typically 5-6 two-fold dilutions are prepared for each compound. Dilutions of compounds are then added to media containing 0.8% CMC resulting in a final 1x drug concentration. After the virus-infected cells have incubated for 1 h at 37°C, the virus innoculum is removed and 1 ml of media containing 0.8% CMC and the various concentrations of compound is added to each well of the dish.

The dish is incubated at 37°C for approximately 2-3 days (HSV-1 and HSV-2) or 6-9 days (HCMV) to allow sufficient growth of virus to form plaques in the cell monolayer. Plaques can be observed and counted microscopically or by staining the cells with 0.1% crystal violet in 20% ethanol. Virus inhibition is determined for each drug concentration by comparing the number of plaques in drug-containing wells to control wells that did not contain drug. Antiviral activity of a compound is expressed as the concentration of compound predicted to reduce the number of plaques in a well by 50% (IC₅₀). The IC₅₀ values are calculated by plotting the per cent inhibition vs. concentration of compound using EXCEL software for linear regression.

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Selection of 4-oxo-DHQ resistant HSV-1 and HSV-2

Vero cells are plated out at a density of 3.5×10^5 cells per well in a six well tissue culture plate. Cells are infected with HSV-1 KOS at a multiplicity of infection (moi) of 0.1 pfu/cell and 1 h post infection the cells are overlayed with 3 ml media containing 20 uM of a 4-oxo-DHQ. Cultures are incubated for 20 h at 37°C, freeze/thawed to release cell-associated virus, and 0.1 ml of culture is used to infect a new monolayer of Vero cells (one passage). Serial passage is repeated seven times in the presence of 20 uM drug. Virus isolates are then plaque purified three times prior to preparation of stocks. Virus recovered from each passage in the presence of compound No. 17 is shown in Figure 3. 4-oxo-DHQ resistant HSV-1 and HSV-2 may also be selected by the marker transfer method described below using wild-type HSV DNA and the corresponding mutant HSV polymerase gene.

Marker Transfer of a HCMV Mutation

A plasmid containing the wild-type HCMV polymerase gene is modified to contain the V823A or V823A and V824L mutations using a site-directed mutagenesis Kit (Stratagene Corp.) and following the manufactures's protocol. HFF cells are plated into T25 tissue culture flasks to achieve 80% confluency at the time of the transfection. Wild type HCMV AD169 DNA and plasmid DNA containing the mutant HCMV polymerase gene are mixed at a ratio of 1:2 (2ug of viral DNA to 4 ug of plasmid DNA). DNA's are transfected using superfect transfection reagent according to methods recommended by the manufacturer (Quiagen Inc.). Cells are harvested five days posttransfection, freeze-thawed to release virus and half of the sample is used to infect HFF cell monolayers. Cells are overlayed with media containing 20 uM 4-oxo-DHQ compound 2 (see Figure 1). Serial

passage is repeated seven times in the presence of 20 uM compound 2 and virus isolates are then plaque purified three times prior to preparation of viral stock.

Isolation of HSV and HCMV viral DNA

HSV DNA is purified from the cytoplasm of infected Vero cells. Vero cells (50 % 5 confluent) are infected at an multiplicity of 0.01 PFU/cell. At 3-5 days postinfection infected cells (100% cpe) are harvested by centrifugation at 1000 rpm in a Beckman GS-6R centrifuge. The pelleted cells are resuspended in TE buffer and placed on ice for 15 minutes. NP-40 is then added to a final concentration of 0.2% and incubated on ice for a further 15 minutes. The cells are centrifuged at 2000 rpm for 10 minutes in a Beckman 10 GS-6R centrifuge. The supernatant is removed and EDTA is added to a final concentration of 20 mM followed by the addition of SDS to a final concentration of 0.3% and proteinase K to a concentration of 50 ug/ml then incubated at 45C for 2 hours. HCMV DNA is isolated by infecting HFF cells (25% confluency) with HCMV at an multiplicity of 0.1 PFU/cell. Cells and media are harvested 5-7 days postinfection (100% cpe) and subjected to low 15 speed centrifugation to remove intact cells and cell debris followed by a high speed spin to pellet virus particles (2500 rpm's in a Beckman SW28 rotor for 1 hour). Following incubation of the HSV and HCMV samples, 1.5 volumes of saturated NaI is added to the digested extract and the refractive index is adjusted to 1.434 -1.435. Ethidium bromide is added to a final concentration of 50 ug/ml. The samples are loaded into a VTI 50centrifuge 20 tube and spun for 24 hours at 45,000 rpm. The DNA band is harvested extracted three times with n-butanol, then dialyzed against TE buffer followed by a dialysis against 95% ethanol and a final dialysis against TE buffer.

DNA Sequencing

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HSV-1, HSV-2 or HCMV viral DNA's are sequenced directly using an ABI377 fluorescence sequencer (Perkin Elmer Applied Biosystems, Foster City, CA) and the ABI BigDye PRISMTM dRhodamine Terminator Cycle Sequencing Ready Reaction Kit with AmpliTaq FSTM DNA polymerase (PE Applied Biosystems). Each cycle sequencing reaction contained about 1.0 ug of purified viral DNA. Cycle-sequencing is performed using an initial denaturation at 98°C for 1 min, followed by 50 cycles: 98°C for 30 sec, annealing at 50°C for 30 sec, and extension at 60°C for 4 min. Temperature cycles and times are controlled by a Perkin-Elmer 9700 thermocycler. Extension products are

purified using CentriflexTM gel filtration cartridges (Edge BioSystems, Gaithersburg, MD). Each reaction product is loaded by pipette onto the column, which is then centrifuged in a swinging bucket centrifuge (Sorvall model RT6000B table top centrifuge) at 750 x g for 1.5 min at room temperature. Column-purified samples are dried under vacuum for about 40 min and then dissolved in 4 ul of a DNA loading solution (83% deionized formamide, 8.3 mM EDTA, and 1.6 mg/ml Blue Dextran). The samples are then heated to 90°C for two min, and held at 4°C until loading. 1.5 ul of each sample is loaded into a single well of the ABI377 sequencer. Sequence chromatogram data files from the ABI377 are analyzed with the computer program Sequencher (Gene Codes, Ann Arbor, MI), for assembly of sequence fragments and correction of ambiguous base calls. Generally sequence reads of 600-700 bp are obtained. Potential sequencing errors are minimized by obtaining sequence information from both DNA strands and by re-sequencing difficult areas using primers at different locations until all sequencing ambiguities are removed.

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The entire coding region of the polymerase genes from both the parent strains and the resistant viruses are sequenced. The DNA sequencing is done using viral DNA as the template thus avoiding cloning of the polymerase genes. The amino acid sequence of the DNA polymerases of HSV-1 KOS, F, Patton and DJL and HSV-2 MS and 186 are compared in Figure 4. Amino acids that are identical for the six polymerases are shaded in black while regions where amino acid differences are found are shaded in gray. The amino acid sequence of the four HSV-1 polymerases are essentially identical with only a few minor changes noted between the different HSV-1 strains. The majority of amino acid changes are found when the sequences of the HSV-1 and HSV-2 polymerases are compared.

Isolation and Characterization of HSV-1 and HSV-2 Mutants That Are Resistant To the 4-oxo-DHQ's and 4-oxo-DHTP Compounds

A panel of viruses consisting of four strains of HSV-1 (KOS, F, DJL, Patton) and three strains of HSV-2 (MS, 35D, 186) are tested in a plaque reduction assay against four different 4-oxo-DHQ compounds (# 1, 2, 4, 5 as shown in Figure 1), and one 4-oxo-DHTP compound (# 3 as shown in Figure 1) and against Acyclovir. The six drugs inhibited replication of the seven virus strains with IC₅₀ values ranging from 2-10 μM (Table 1). In order to select for 4-oxo-DHQ resistant mutants, HSV-1 strains KOS, F, and DJL along with HSV-2 strains 186 and MS are serially passaged in the presence of 20 μM compound

1. Following the seventh passage, 4-oxo-DHQ resistant virus from each strain are plaque purified three times and high-titer stocks are made. All of the resistant HSV mutants grew to high titers in Vero cells, indicating that the mutations in the resistant isolates did not significantly impair their growth. The mutants selected with 4-oxo-DHQ compound 1 exhibited >10 fold increase in IC₅₀ when tested in a plaque reduction assay against 4-oxo-DHQ compound 1 Data are shown in Table 2.

Table 2
4-oxo-DHQ Resistant Virus of HSV-1 and HSV-2

Virus Mutants	Compound 1 IC ₅₀ (uM)	Amino Acid Change in HSV DNA Polymerase
HSV-1 Kos-M1	>20	- V823A
HSV-1 F-M1	>20	- V823A
HSV-1 DJL-M1	>20	-V823A
HSV-2 MS-M1	>20	- V826A
HSV-2 186-M1	>20	- V828A

*HSV-1 and HSV-2 isolates grown in the presence of 4-oxo-DHQ select for resistant virus.

DNA sequence analysis of the 4-oxo-DHQ resistant mutants (HSV-1 KOS-M1, HSV-1 F-M1, HSV-1 DJL-M1, HSV-2 186-M1, HSV-2 MS-M1) demonstrated that all five mutants contained a single point mutation of T to C at the binding domain resulting in a Valine to Alanine amino acid change.

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<u>Isolation and Characterization of A HCMV Mutant That Is Resistant to The 4-oxo-DHQ's and 4-oxo-DHTP Compounds</u>

In order to select for a 4-oxo-DHQ HCMV resistant mutant, virus (strain AD169) is serially passaged in the presence of 20 uM a 4-oxo-DHQ. Although we could readily select for HSV mutants using this procedure we failed to isolate an HCMV mutant, even when the virus is passaged at low drug concentrations (<5 uM). Comparison of the amino acid sequence of the HSV polymerase, Y-G-F-T-G-V-Q-H-G, and HCMV polymerase, Y-G-F-T-G-V-V-N-G, in the region of amino acid 823 (underlined amino acid) showed that there is a second valine at position 824 in the HCMV polymerase. In order to determine if both valines need to be changed in order to confer resistance to the 4-oxo-DHQ's, *in vitro* polymerase assays are done using mutant HCMV polymerases containing either V823A or V823A plus V824L (Table 3).

Table 3

HCMV Mutant Polymerase Exhibits Resistance to 4-oxo-DHQ*

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Polymerase	Compound 1 IC ₅₀ (uM)
HCMV (wild)	4.6
HCMV V823A	17.2
HCMV V823A/V824L	42.9

^{*}Generation of the valine to alanine at amino acid 823 of HCMV results in a 3.5-fold increase in resistance.

The V823A alone resulted in a 3.5-fold increase in the IC₅₀ while the polymerase with the double amino acid change had nearly 10-fold increase in the IC₅₀. In order to isolate an HCMV resistant mutant marker rescue experiments are done. Plasmids containing the mutant polymerase genes are transfected into HFF cells along with wild type HCMV AD169 DNA. The resulting virus is then serially passaged in the presence of 20 uM compound 1 (see figure 1). A 4-oxo-DHQ resistant virus is isolated from marker rescue studies done with the HCMV polymerase gene containing mutations that result in the V823A, V824L amino acid changes, but not with the gene containing V823A change alone. The mutant selected with compound 1 (HCMV AD169-M1) exhibited ~7-fold increase in IC₅₀ when tested in a plaque reduction assay compared to Ganciclovir and cidofovir which has a \leq 2-fold change in sensitivity (Table 4).

Table 4
Plaque reduction assay of 4-oxo-DHQ resistant HCMV*

Drug	HCMV AD169 IC ₅₀ (μM)	HCMV AD169 – M1 IC ₅₀ (μM)		
Compound 1	0.7	4.7		
Ganciclovir	0.9	1.0		
Cidofovir	0.3	0.6		

^{*}Recombination of wild-type HCMV with a polymerase gene containing the valine to alanine at amino acid 823 and the valine to leucine at amino acid 824 allowed for selection of resistant virus with about 7-fold less sensitivity to compound 1.

^{*}Mutation of the amino acid from valine to alanine and amino acid 824 from valine to leucine results in an 9-fold increase in resistance, relative to wild type.

^{*}Sensitivity of resistant HCMV virus to Ganciclovir and Cidofovir verifies that the 4-oxo-DHQ's mechanism for inhibiting the polymerase protein is unique

The entire coding region of the HCMV polymerase genes from both the parent strain and the resistant virus are sequenced. The DNA sequencing is again done using viral DNA as the template thus avoiding cloning of the polymerase genes. Comparison of the DNA sequence of the two polymerase genes demonstrated that the resistant mutant contained two point mutations that resulted in the predicted V823A, V824L amino acid changes. As with the HSV resistant viruses these results demonstrate the critical role of the region encompassing amino acid 823 for inhibition of polymerase activity by these compounds.

10 Antiviral Activity of Nucleoside and Non-Nucleoside Polymerase Inhibitors Against 4oxo-DHQ Resistant Mutants

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In order to determine if the 4-HQ binding domain mutations alter the sensitivity of the HSV-1, HSV-2 and HCMV mutants to both non-nucleoside (4-oxo-DHQ's) and nucleoside inhibitors (e.g Acyclovir and ganciclovir) several of the mutants are tested in plaque reduction assays against a series of non-nucleoside compounds including Foscarnet (PFA), 4-HQ's 4-oxo-DHQ's and 4-oxo-DHTP's (Table 5). The mutants are also tested against a series of nucleoside inhibitors including acyclovir and ganciclovir (Table 5). The activity of these compounds against the mutants is compared to their activity against the wild type strains that are used to isolate the HSV and HCMV mutants. When tested against a number of 4-HQ's, 4-oxo-DHQ's and 4-oxo-DHTP's and other related classes of compounds all of the drugs are found to inhibit the wild type virus with IC50 values ranging from <0.1 uM to 30 uM. When these drugs are tested against the resistant viruses they are found to have IC₅₀ values 5 to 10 fold higher then the parent virus. There is little if any difference in the IC₅₀ values of the nucleoside compounds and the non-nucleoside PFA between the wild type and mutant HSV-1, HSV-2, and HCMV viruses. These results demonstrate that the amino acid change in the binding domain (V823A in the HSV-1 polymerase, V826A in the HSV2-MS polymerase, V828A in the HSV2-186 polymerase, and the V823A/V824L changes in the HCMV polymerase) resulted in resistance to the 4oxo-DHQ's and 4-oxo-DHTP's, which provides further evidence that these classes of compounds share an affinity for a region we refer to as the binding domain. In contrast, these amino acid changes did not alter the activity of these viruses to other classes of polymerase inhibitors.

Table 5

Antiviral activity of nucleoside and non-nucleoside polymerase inhibitors against HSV-1, HSV-2, and HCMV Isolates selected for 4-oxo-DHQ resistance*

	Plaque Reduction Assay – IC ₅₀ (μM)									
Drug	HSV-2 MS	HSV-2 MS-M1	HSV-1 KOS	HSV-1 KOS-M1	HCMV AD169	HCMV AD169-M1				
6	28.8	>50	24.6	>50	5.1	>16				
7	8.8	27.9	6.5	>50	0.3	3.4				
8	2.3	>50	5.1	>50	< 0.1	1.1				
9	0.9	48.7	1.9	>50	<0.1	3.1				
10	29.2	>50	15.8	>50	1.1	>16				
11	3.0	>50	3.1	>50	0.7	3.9				
12	0.4	12.5	1.3	>50	0.2	1.1				
13	5.3	>50	5.5	<25	2.7	>16				
14	1.6	>50	28.4	>50	0.9	18.4				
2	1.3	>50	3.3	>50	0.4	4.0				
4	2.1	28.4	4.2	>50	0.6	2.1				
3	0.8	>50	4.0	>50	1.5	6.2				
15	5.9	>50	>50	>50	0.7	7.7				
Tudr	5.0	6.1	1.1	0.8	ND	ND				
Bvdu	5.8	5.9 ·	2.1	0.1	ND	ND				
ACV	2.4	2.8	3.9	4.4	ND	ND				
AraC	0.2	0.1	0.2	0.2	ND	ND				
AraT	6.6	3.6	11.6	3.6	ND	ND				
AraA	10.6	18.2	26.1	27.2	ND	ND				
GCVir	ND	ND	ND	ND	0.8	0.8				
CDV	ND	ND	ND	ND	0.4	0.3				
PFA	ND	ND	ND	ND	38	<20				

^{*}HSV-2 MS, HSV-1 KOS, HCMV AD169: wild type strains

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Antiviral compounds identified by the present invention can conveniently be administered in a pharmaceutical composition containing the compound in combination with a suitable excipient, the composition being useful in combating viral infections. Pharmaceutical compositions containing a compound appropriate for antiviral use are prepared by methods and contain excipients which are well known in the art. A generally recognized compendium of such methods and ingredients is Remington's Pharmaceutical Sciences by E.W. Martin (Mark Publ. Co., 15th Ed., 1975).

Antiviral compounds identified by the present invention and their compositions can be administered parenterally (for example, by intravenous, intraperitoneal or intramuscular

^{*}HSV-2 MS-M1, HSV-1 KOS-M1, HCMV AD169-M1: mutants selected for 4-oxo-DHQ resistance *ND - Not Done.

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injection), topically, orally, or rectally, depending on whether the preparation is used to treat internal or external viral infections.

For oral therapeutic administration, the active compound may be combined with one or more excipients and used in the form of ingestible tablets, buccal tablets, troches, capsules, elixirs, suspensions, syrups, wafers, and the like. Such compositions and preparations should contain at least 0.1% of active compound. The percentage of the compositions and preparations may, of course, be varied and may conveniently be between about 2 to about 60% of the weight of a given unit dosage form. The amount of active compound in such therapeutically useful compositions is such that an effective dosage level will be obtained.

The tablets, troches, pills, capsules, and the like may also contain the following: binders such as gum tragacanth, acacia, corn starch or gelatin; excipients such as dicalcium phosphate; a disintegrating agent such as corn starch, potato starch, alginic acid and the like; a lubricant such as magnesium stearate; and a sweetening agent such as sucrose, fructose, lactose or aspartame or a flavoring agent such as peppermint, oil of wintergreen, or cherry flavoring may be added. When the unit dosage form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier, such as a vegetable oil or a polyethylene glycol. Various other materials may be present as coatings or to otherwise modify the physical form of the solid unit dosage form. For instance, tablets, pills, or capsules may be coated with gelatin, wax, shellac or sugar and the like. A syrup or elixir may contain the active compound, sucrose or fructose as a sweetening agent, methyl and propylparabens as preservatives, a dye and flavoring such as cherry or orange flavor. Of course, any material used in preparing any unit dosage form should be pharmaceutically acceptable and substantially non-toxic in the amounts employed. In addition, the active compound may be incorporated into sustained-release preparations and devices.

Antiviral compounds identified by the present invention and their compositions can also be administered intravenously or intraperitoneally by infusion or injection. Solutions of the active compound or its salts can be prepared in water, optionally mixed with a nontoxic surfactant. Dispersions can also be prepared in glycerol, liquid polyethylene glycols, triacetin, and mixtures thereof and in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

Pharmaceutical dosage forms suitable for injection or infusion can include sterile aqueous solutions or dispersions or sterile powders comprising the active ingredient which

are adapted for the extemporaneous preparation of sterile injectable or infusible solutions or dispersions, optionally encapsulated in liposomes. In all cases, the ultimate dosage form should be sterile, fluid and stable under the conditions of manufacture and storage. The liquid carrier or vehicle can be a solvent or liquid dispersion medium comprising, for example, water, ethanol, a polyol (for example, glycerol, propylene glycol, liquid polyethylene glycols, and the like), vegetable oils, nontoxic glyceryl esters, and suitable mixtures thereof. The proper fluidity can be maintained, for example, by the formation of liposomes, by the maintenance of the required particle size in the case of dispersions or by the use of surfactants. The prevention of the action of microorganisms can be brought about by various antibacterial and antifungal agents, for example, parabens, chlorobutanol, phenol, sorbic acid, thimerosal, and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars, buffers or sodium chloride. Prolonged absorption of the injectable compositions can be brought about by the use in the compositions of agents delaying absorption, for example, aluminum monostearate and gelatin.

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Sterile injectable solutions can be prepared by incorporating the active compound in the required amount in the appropriate solvent with various of the other ingredients enumerated above, as required, followed by filter sterilization. In the case of sterile powders for the preparation of sterile injectable solutions, the preferred methods of preparation are vacuum drying and the freeze drying techniques, which yield a powder of the active ingredient plus any additional desired ingredient present in the previously sterile-filtered solutions.

For topical administration, the present compounds may be applied in pure form, i.e., when they are liquids. However, it will generally be desirable to administer them to the skin as compositions or formulations, in combination with a dermatologically acceptable carrier, which may be a solid or a liquid.

Useful solid carriers include finely divided solids such as talc, clay, microcrystalline cellulose, silica, alumina and the like. Useful liquid carriers include water, alcohols or glycols or water-alcohol/glycol blends, in which the present compounds can be dissolved or dispersed at effective levels, optionally with the aid of non-toxic surfactants. Adjuvants such as fragrances and additional antimicrobial agents can be added to optimize the properties for a given use. The resultant liquid compositions can be applied from absorbent pads, used to impregnate bandages and other dressings, or sprayed onto the affected area using pump-type or aerosol sprayers. Thickeners such as synthetic polymers, fatty acids,

fatty acid salts and esters, fatty alcohols, modified celluloses or modified mineral materials can also be employed with liquid carriers to form spreadable pastes, gels, ointments, soaps, and the like, for application directly to the skin of the user.

Examples of useful dermatological compositions which can be used to deliver the compounds of formula I to the skin are known to the art; for example, see Jacquet et al. (U.S. Pat. No. 4,608,392), Geria (U.S. Pat. No. 4,992,478), Smith et al. (U.S. Pat. No. 4,559,157) and Wortzman (U.S. Pat. No. 4,820,508).

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Useful dosages of the compounds of formula I can be determined by comparing their *in vitro* activity, and *in vivo* activity in animal models. Methods for the extrapolation of effective dosages in mice, and other animals, to humans are known to the art; for example, see U.S. Pat. No. 4,938,949.

The compound is conveniently administered in unit dosage form; for example, containing 5 to 1000 mg, conveniently 10 to 750 mg, most conveniently, 50 to 500 mg of active ingredient per unit dosage form. The desired dose may conveniently be presented in a single dose or as divided doses administered at appropriate intervals, for example, as two, three, four or more sub-doses per day. The sub-dose itself may be further divided, e.g., into a number of discrete loosely spaced administrations; such as multiple inhalations from an insufflator or by application of a plurality of drops into the eye.

For internal infections, the compositions can be administered orally or parenterally at dose levels, calculated as the free base, of about 0.1 to 300 mg/kg, preferably 1.0 to 30 mg/kg of mammal body weight, and can be used in man in a unit dosage form, administered one to four times daily in the amount of 1 to 1000 mg per unit dose.

For parenteral administration or for administration as drops, as for eye infections, the compounds are presented in aqueous solution in a concentration of from about 0.1 to about 10%, more preferably about 0.1 to about 7%. The solution may contain other ingredients, such as emulsifiers, antioxidants or buffers.

Generally, the concentration of the compound(s) of formula I in a liquid composition, such as a lotion, will be from about 0.1-25 wt-%, preferably from about 0.5-10 wt-%. The concentration in a semi-solid or solid composition such as a gel or a powder will be about 0.1-5 wt-%, preferably about 0.5-2.5 wt-%.

The exact regimen for administration of the compounds and compositions disclosed herein will necessarily be dependent upon the needs of the individual subject being treated, the type of treatment and, of course, the judgment of the attending practitioner.

The antiviral activity of a compound of the invention can be determined using pharmacological models which are well known to the art, or using Test A described below.

The compounds of formula (I) and pharmaceutically acceptable salts thereof are useful as antiviral agents. Thus, they are useful to combat viral infections in animals, including man. The compounds are generally active against herpes viruses, and are particularly useful against the varicella zoster virus, the Epstein-Barr virus, the herpes simplex virus, the human herpes virus type 8 (HHV-8) and the cytomegalovirus (CMV).

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CLAIMS

We claim:

1. A method of selecting compounds that inhibit herpes viruses comprising:

- a) measuring IC₅₀ of a compound of interest that inhibits a wild type herpes virus,
- 5 b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant herpes virus which is the same strain as the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - selecting the compound of interest wherein the IC_{50} of step b is at least 3 times greater than the IC_{50} of step a.

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- 2. A method of selecting compounds that inhibit herpes viruses comprising:
- a) measuring IC₅₀ of a compound of interest that inhibits a binding domain mutant herpes virus,
- b) measuring IC_{50} of the same compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes virus,
- c) comparing IC₅₀ of step a with IC₅₀ of step b; and
- d) selecting the compound of interest wherein the IC_{50} of step a is at least 3 times greater than the IC_{50} of step b.
- The method of claim 1 or 2 wherein the herpes virus is HSV-1, HSV-2, HCMV, VZV, EBV, or HHV-8.
 - 4. A method of selecting compounds that inhibit herpes viruses comprising:
 - a) measuring IC₅₀ of a compound of interest that inhibits a wild type HSV-1,
- 25 b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant HSV-1 which is the same strain as the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - d) selecting the compound of interest wherein the IC_{50} of step b is at least 3 times greater than the IC_{50} of step a.

- 5. A method of selecting compounds that inhibit herpes viruses comprising:
- a) measuring IC₅₀ of a compound of interest that inhibits a binding domain mutant HSV-1,

b) measuring IC_{50} of the same compound that inhibits a wild type herpes virus which is the same strain as the mutant HSV-1,

- c) comparing IC₅₀ of step a with IC₅₀ of step b; and
- d) selecting the compound of interest wherein the IC₅₀ of step a is at least 3 times greater than the IC₅₀ of step b.
 - 6. The method of claim 4 or 5 wherein HSV-1 is HSV-1 KOS, HSV-1 F, HSV-1 DJL or HSV-1 Patton.
- 7. The method of claim 5 or 6 wherein the mutation of a wild type herpes virus to mutant herpes virus is at amino acid 823 from valine to alanine.
 - 8. A method of selecting compounds that inhibit herpes viruses comprising:
 - a) measuring IC₅₀ of a compound of interest that inhibits a wild type HSV-2,
- measuring IC₅₀ of the same compound that inhibits a binding domain mutant HSV-2 which is the same strain as the wild type herpes virus,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - d) selecting the compound of interest wherein the IC₅₀ of step b is at least 3 times greater than the IC₅₀ of step a.

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- 9. A method of selecting compounds that inhibit herpes viruses comprising:
- a) measuring IC₅₀ of a compound of interest that inhibits a binding domain mutant HSV-2,
- b) measuring IC₅₀ of the same compound that inhibits a wild type herpes virus which is the same strain as the mutant HSV-2,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and
 - d) selecting the compound of interest wherein the IC_{50} of step a is at least 3 times greater than the IC_{50} of step b.
- The method of claim 8 or 9 wherein HSV-2 is HSV-2 MS, HSV-2 35D, or HSV-2 186.
 - 11. A method of selecting compounds that inhibit herpes viruses comprising:

a) measuring IC₅₀ of a compound of interest that inhibits a wild type HCMV,

- b) measuring IC₅₀ of the same compound that inhibits a binding domain mutant HCMV which is the same strain as the wild type herpes virus,
- c) comparing IC₅₀ of step a with IC₅₀ of step b; and
- selecting the compound of interest wherein the IC₅₀ of step b is at least 3 times greater than the IC₅₀ of step a.
 - 12. A method of selecting compounds that inhibit herpes viruses comprising:
 - a) measuring IC₅₀ of a compound of interest that inhibits a binding domain mutant HCMV,
 - b) measuring IC₅₀ of the same compound that inhibits a wild type herpes virus which is the same strain of the mutant HCMV,
 - c) comparing IC₅₀ of step a with IC₅₀ of step b; and

- d) selecting the compound of interest wherein the IC₅₀ of step a is at least 3 times greater than the IC₅₀ of step b.
 - 13. The method of claim 8 or 9 wherein HCMV is AD169.
- 14. The methods of claims 1, 4, 8, or 11 wherein IC₅₀ of step b is at least 5 times greater than the IC₅₀ of step a.
 - 15. The methods of claims 2, 5, 9, or 12 wherein IC_{50} of step a is at least 5 times greater than the IC_{50} of step b.
- 25 16. A use of compounds for manufacturing of medicinals for selectively treating diseases caused by herpes viruses in a human host comprising administering a compound to a human in need of such treatment wherein said compound inhibits herpes viruses by interaction with the binding domain in the viral DNA polymerase.
- 17. A use of compounds for manufacturing of medicinals for selectively inhibiting herpes viruses in a human host comprising administering a compound to a human in need of such treatment wherein IC₅₀ of the compound that inhibits a binding domain

mutant herpes virus is at lease 3 times greater than IC₅₀ of the compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes virus.

- 18. The use of claim 17 wherein IC₅₀ of the compound that inhibits a binding domain mutant herpes virus is at lease 5 times greater than IC₅₀ of the compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes viruse.
- 19. The use of claim 17 wherein herpes viruses is HSV-1, HSV-2, HCMV, VZV, EBV, or HHV-8.
 - 20. A use of compounds for manufacturing of medicinals for treating herpesviral infections in a human host wherein IC₅₀ of the compound that inhibits a binding domain mutant herpes virus is at lease 5 times greater than IC₅₀ of the compound that inhibits a wild type herpes virus which is the same strain as the mutant herpes virus.

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- 21. A use of compounds for manufacturing of medicinals for treating herpesviral infections in a human host wherein said compound inhibits the herpesvirus by interacting with the binding domain in the viral DNA polymerase.
- 22. The herpesviral infection of claim 20 or 21 which is HSV-1, HSV-2, HCMV, VZV, EBV, or HHV-8 infection.
- 23. A compound for the inhibiting of herpesvirus DNA polymerases wherein passage of a wild type herpes virus in the presence of said compound results a change of the wild type HSV-1 polymerases at amino acid 823 from valine to alanine.
- 24. A compound for inhibiting herpesvirus DNA polymerases wherein passage of a wild type herpes virus in the presence of said compound results in a change of the wild type HCMV polymerases at amino acid 823 from valine to alanine and at amino acid 824 from valine to leuline.

25. A mutant herpesvirus DNA molecule having a nucleotide sequence selected from a group consisting of SEQ.ID.NO. 1; SEQ.ID.NO. 3; SEQ.ID.NO. 5; SEQ.ID.NO. 7; SEQ.ID.NO. 9; and SEQ.ID.NO. 11.

5 26. A mutant herpesvirus polymerase amino acid molecule having an amino acid sequence selected from a group consisting of SEQ.ID.NO. 2; SEQ.ID.NO. 4; SEQ.ID.NO. 6; SEQ.ID.NO. 8; SEQ.ID.NO. 10 and SEQ.ID.NO. 12.

Figure 1 4-HQ, 4-oxo-DHQ and 4-oxo-DHTP antiviral compounds

(Figure 1 continue)

Compound No. 7

(Figure 1 continue)

(Figure 1 continue)

Compound No.15

Compound 17

Figure 2. The HSV1 (KOS Strain) DNA Polymerase Amino Acid 823 is Critical for Resistance to 4-Hydroxyquinolines and Related Compounds

NH-	A	11	VI	111	-	-		v I	Нсоон
				V82	3A				
HS/	/1-KOS-M1 Y	G	F '	T G	À	Q	н	G - 826	
HS/		G	F '	T G	٧	Q	Н	G - 826	
HS\		G	F .	T G	V	Q	Н	G - 829	
VZV		G	F '	T G	٧	A	Q	G - 791	
EB!		G	F '	T G	٧	A	N	G - 696	
HC		G	F	TG	٧	٧	N	G - 826	
HH		G	V '	T G	Α	A	Н	G - 681	
HH		G	٧ .	T G	Α	T	Н	S - 681	
HH		G	F	T G	V	A	S	G - 696	

Schematic of HSV1 polymerase illustrating the conserved regions A and I-VI found in class 2 polymerases. Also shown are the amino acid sequence for the highly conserved herpesvirus domain in region III which surrounds the HSV1 amino acid 823.

Figure 3 Serial Passage of HSV-1 in Presence of 20 μM compound 17

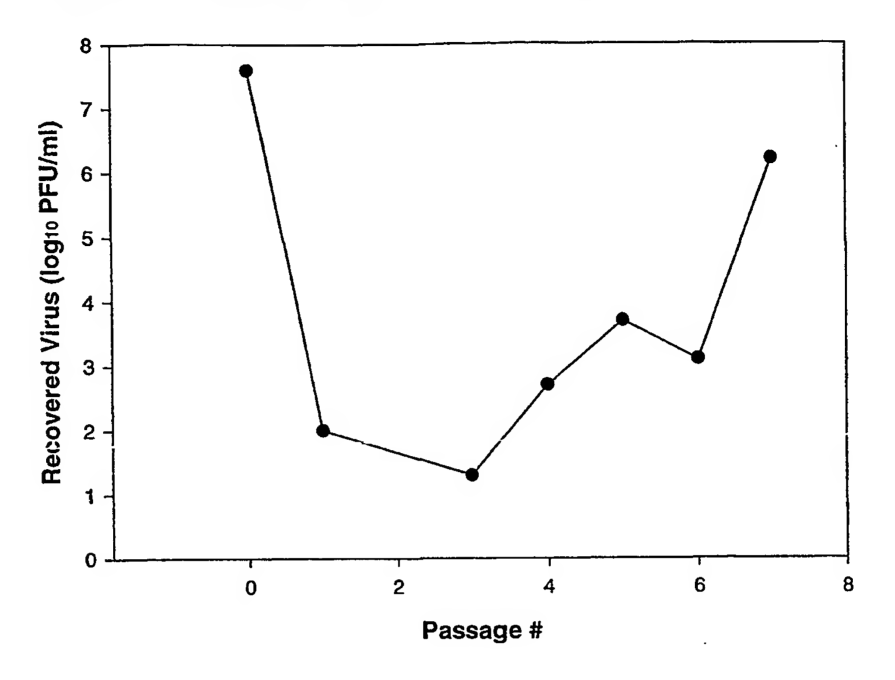


Figure 4 Comparison of Wild type HSV-1 and HSV-2 DNA Polymerases Amino Acid Sequences Alligned by Amino Acid Homology*

		B					•
	HSV2-MS	MFCAAGGPTS	PGGKSAARAA	SGFFAPHNPR	GATOTAPPPC	RRONFYNPHL	-50
						RRONFYNPHL	
	HSV2-186	•	PGGKSAARAA				
5	HSV1-Kos	MFSGGGGPLS	PGGKSAARAA			LRQNFYNPYL	
	HSV1-Patton	MFSGGGGPLS	PGGKSAARAA	SGFFAPAGPR	GAGR.GPPPC	LRONFYNPYL	-49
			PGGKSAARAA			LRQNFYNPYL	-49
	HSV1-DJL						
	HSV1-F	MFSGGGGPLS	PGGKSAARAA	SGFFAPAGPR	GAGR.GPPPC	LRQNFYNPYL	-43
10	HSV2-MS	AQTGTQPKAP	GPAORHTYYS	ECDEFRFIAP	RSLDEDAPAE	QRTGVHDGRL	-100
	HSV2-186	AOTGTOPKAP		ECDEFRETAP	RSLDEDAPAE	QRTGVHDGRL	-100
		~					-99
	HSV1-Kos	APVGTQQKPT		ECDEFRFIAP			• -
	HSV1-Patton	APVGTQQKPT	GPTQRHTYYS	ECDEFRFIAP	RVLDEDAPPE		-99
	HSV1-DJ1	APVGTQQKPT	GPTORHTYYS	ECDEFRFIAP	RVLDEDAPPE	KRAGVHDGHL	-99
15		APVGTQQKPT		ECDEFRFIAP	RVLDEDAPPE	KRAGVHDGHL	-99
15	HSV1-F	APVGIQQRFI	GETÖKUTTIS	ECDELIG XXX		144101111111111111111111111111111111111	
				naminani at	LICCADITA DEC	FDPTVTVFHV	-150
	HSV2-MS	RRAPKVYCGG	DERDVLRVGP	EGFWPRRLRL			
	HSV2-186	PRAPKVYCGG	DERDVLRVGP	EGFWPRRLRL	WGGADHAPEG	FDPTVTVFHV	-15û
	HSV-Kos	KRAPKVYCGG	DERDVLRVGS	GGFWPRRSRL	WGGVDHAPAG	FNPTVTVFHV	-149
00				GGFWPRRSRL		FNPTVTVFHV	-149
20	HSV1-Patton	KRAPKVYCGG	DERDVLRVGS				
	HSV1-DJL	KRAPKVYCGG	DERDVLRVGS	GGFWPRRSRL		FNPTVTVFHV	-149
	HSV1-F	KRAPKVYCGG	DERDVLRVGS	GGFWPRRSRL	WGGVDHAPAG	FNPTVTVFHV	-149
	*******	SENTE DUNDUM	VCMD A A OT UE	RFMDAITPAG	TVITLLGLTP	EGHRVAVHVY	-200
	HSV2-MS		YSMRAAQLHE				
25	HSV2-186	ADIPEHAEHY	YSMRAAQLHE	RFMDAITPAG		EGHRVAVHVY	-200
	HSV-Kos	YDILENVEHA	YGMRAAQFHA	RFMDAITPTG	TVITLLGLTP	EGHRVAVHVY	
	HSV1-Patton	VIDTI.ENVEHA	YCMRAAOFHA	REMDATTETG	TVITLLGLTP	EGHRVAVHVY	-199
		TOTELLIA A DITT	TOTAL A VISITA	שתחת לחושים כי	TVITLLGLTP	EGHRVAVHVY	_199
	HSV1-DJL						
	HSV1-F	YDILENVEHA	YGMRAAQFHA	RFMDAITPTG	TVITLLGLTP	EGHRVAVHVY	-199
30							
50	HSV2-MS	CUIDOALAMVIK	AEVDRHI.OCR	APRDLCERLA	AALRESPGAS	FRGISADHFE	-250
		GINGILIIM	ABVDIUIDQCIC	A DODE CERT A	AATDECDCAC	FRGISADHFE	-250
	HSV2-186	GTRQYFYMNK	AEVDRHLQCK	APROLCERLA	AALRESPGAS	PRGISADHFE	_
	HSV-Kos	GTRQYFYMNK	EEVDRHLQCR	APRDLCERMA	AALRESPGAS	FRGISADHFE	
	HSV1-Patton	GTROYFYMNK	EEVDRHLOCR	APRDLCERMA	AALRESPGAS	FRGISADHFE	-249
25			EETDDUI OCD	A DDDI CERMA	AALRESPGAS	FRGISADHFE	-249
35	HSV1-DJL	GTRQIF IMM	EEADKUDÕCK	APRODUCERUM			-249
	HSV1-F	GTRQYFYMNK	EEVDRHLQCR	APRDLCERMA	AALRESPGAS	FRGISADHFE	-243
							000
	HSV2-MS	AEVVERADVY	YYETRPTLYY	RVFVRSGRAL	AYLCDNFCPA	IRKYEGGVDA	-300
	HSV2-186	AEVVERADVY	YYETRPTLYY	RVFVRSGRAL	AYLCDNFCPA	IRKYEGGVDA	-300
40		EVVERTDVY Y					-299
40							
	HSV1-Patton					IKKYEGGVDA	-
	HSV1-DJL				SYLCDNFCPA		-299
	HSV1-F	AEVVERTDVY	YYETRPALFY	RVYVRSGRVL	SYLCDNFCPA	IKKYEGGVDA	-299
	11571 1	••••					
15	· UCTIO MC	THE TELEVISION OF THE PROPERTY	EUTECMVDI.K	PGRGNAPAQP	RPPTAFGTSS	DVEFNCTADN	-350
45	HSV2-MS						
	HSV2-186			PGRGNAPAQF			
	HSV-Kos	TTRFILDNPG	FVTFGWYRLK	PGRNNTLAQF	RAPMAFGTSS	DVEFNCTADN	-349
	HSV1-Patton	TTRETT, DNPC	FUTEGWYRLK	PGRNNTLAOP	RAPMAFGTSS	DVEFNCTADN	-349
							-349
	HSV1-DJL	T.T.K.E.T.DINE.	FOTEGMIKTY	PGRNNTLAQE		•	
50	HSV1-F	TTRFILDNPG	FVTFGWYRLK	PGRNNTLAQE	RAPMAFGTSS	DVEFNCTADN	-349
	HSV2-MS	LAVECAMODIA	PAYKIMCFDI	ECKAGGEDEL	AFPVAERPED	LVIQISCLLY	-400
		LAVEGAMCDL			AFPVAERPED		-400
	HSV2-186						
	HSV-Kos	LAIEGGMSDL			. AFPVAGHPED		
55	HSV1-Patton	LAIEGGMSDL	PAYKLMCFDI	ECKAGGEDEI	J AFPVAGHPED	LVIQISCLLY	-399
	HSV1-DJL	LAIEGGMSDL		ECKAGGEDET	AFPVAGHPEL	LVIQISCLLY	-399
	HSV1-F	LAIEGGMSDL	PAYKLMCFDI	. ECKAGGEDEL	AFPVAGHPED	, MATOCODI	-333
							4 = 4
	HSV2-MS	DLSTTALEHI	LLFSLGSCDI	PESHLSDLAS	RGLPAPVVLE	FDSEFEMLLA	-450
60	HSV2-186	דים אות און בים	T.T.F.CT.CCCDT	PESHLEDI.A	RGI PAPINTI	FDSEFEMLLA	-450
60							-449
	HSV-Kos				A RGLPTPVVLE		
	HSV1-Patton	DLSTTALEHV	LLFSLGSCDI	PESHLNELA!	A RGLPTPVVLE		
	HSV1-DJL	DLSTTALEHV	LLFSLGSCDI	PESHLNELA	A RGLPTPVVLE	FDSEFEMLLA	-449
	HSV1-F					FDSEFEMLLA	-449
~~	TO A T_L	THETTERN					-
65							

	HSV2-MS	FMTFVKQYGP					
	HSV2-186	FMTFVKQYGP	MT 4 T G T T 1 T T T T T T T T T T T T T T T		TEIYKVPLDG		-500
	HSV-Kos	FMTLVKQYGP	EFVTGYNIIN	FDWPFLLAKL	TDIYKVPLDG		-499
	HSV1-Patton	FMTLVKQYGP	D2	FDWPFLLAKL	TDIYKVPLDG		- 499
5	HSV1-DJL	FMTLVKQYGP	EFVTGYNIIN	FDWPFLLAKL	TDIYKVPLDG		-499
	HSV1-F	FMTLVKQYGP	EFVTGYNIIN	FDWPFLLAKL	TDIYKVPLDG	YGRMNGRGVF	-499
						T NIN 17N TON 17T 77	EEA
	HSV2-MS	_			TDKVKLSSYK TDKVKLSSYK	T NATATATATI Y	-550 -550
	HSV2-186		QKRSKIKVNG		_		-549
10	HSV-Kos	RVWDIGQSHF	QKRSKIKVNG	MVNIDMYGII	TDKIKLSSYK	PNAVAEAVIA	
	HSV1-Patton	RVWDIGQSHF	QKRSKIKVNG	MVNIDMYGII	TDKIKLSSYK	LNAVAEAVLK	
	HSV1-DJL	RVWDIGQSHF	QKRSKIKVNG	MVNIDMYGII	TDKIKLSSYK	LNAVAEAVLK	-549
	HSV1-F	RVWDIGQSHF	QKRSKIKVNG	MVNIDMYGII	TDKIKLSSYK	LNAVAEAVLK	-549
15	HSV2-MS	DKKKDLSYRD	TPAYVASGPA	ORGVIGEYCV	QDSLLVGQLF	FKFLPHLELS	-600
1.5	HSV2-186				QDSLLVGQLF		-600
	HSV-Kos				QDSLLVGQLF		-599
					QDSLLVGQLF		-599
	HSV1-Patton.				QDSLLVGQLF		-599
20	HSV1-DJL				QDSLLVGQLF		-599
20	HSV1-F	DKKKDLSYRD	Irailaaura	5KG A TGETCT	ODPHI AGONI	PREBLIBBIO	
	HSV2-MS	AVARLAGINI	TRTIYDGQQI	RVFTCLLRLA	GQKGFILPDT	QGRFRGLDKE	-650
	HSV2-186				GQKGFILPDT		-650
	HSV-Kos	AVARLAGINI	TRTIYDGQQI	RVFTCLLRLA	DQKGFILPDT	QGRFRGAGGE	-649
25	HSV1-Patton	AVARLAGINI	TRTIYDGQQI	RVFTCLLRLA	DQKGFILPDT	QGRFRGAGGE	-649
	HSV1-DJL	AVARLAGINI	TRTIYDGQQI	RVFTCLLRLA	DQKGFILPDT	QGRFRGAGGE	-649
	HSV1-F				DQKGFILPDT		-649
							605
	HSV2-MS	APKRPAVPRG		DEDKDDDE		VARETGGRHV	
30	HSV2-186	APKRPAVPRG		DEDKDDDEDG			-697
	HSV-Kos	APKRPAAARE			EEGGGEREPE		-694
	HSV1-Patton	APKRPAAARE	DEERP	EEEGEDEDER	EEGGGEREPE		-694
	HSV1-DJL	APKRPAAARE	= =	EEEGEDENER			-694
	HSV1-F	APKRPAAARE	DEERP	EEEGEDEDER	EEGGGEREPE	GARETAGRHV	-694
35				•			
	HSV2-MS	GYQGARVLDP	TSGFHVDPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRPEAVA	-747
	HSV2-186	GYQGARVLDP	TSGFHVDPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRPEAVA	-749
	HSV-Kos	GYQGARVLDP	TSGFHVNPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRADAVA	-744
	HSV1-Patton	GYQGARVLDP	ISGFHVNPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRADAVA	-744
40	HSV1-DJL	GYQGARVLDP	TSGFHVNPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRADAVA	-744
	HSV1-F	GYQGARVLDP	TSGFHVNPVV	VFDFASLYPS	IIQAHNLCFS	TLSLRADAVA	-744
							707
	HSV2-MS	HLEADRDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	-/9/
	HSV2-186	HLEADRDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	- /99
45	HSV-Kos	HLEAGKDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	- 794
	HSV1-Patton	HLEAGKDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	- /94
	HSV1-DJL	HLEAGKDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	-794
	HSV1-F	HLEAGKDYLE	IEVGGRRLFF	VKAHVRESLL	SILLRDWLAM	RKQIRSRIPQ	-794
60	770110 MG	content to	VOON N TVANIC	NCVCETCV	HGLLPCLHVA	ATTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	-847
50	HSV2-MS	STREEWAND	MOOVY TRANSC	MOUVEFOCUO	HGLLPCLHVA	ATUTTTGREM	-849
	HSV2-186	SPPEEAVELD	MOONTHAIL A	MEANGEMEAN	HGLLPCLHVA	ATTITITOREM	-844
	HSV-Kos	SSPEEAVLLD	MOON A TRAMIC	MOVIGE TOVE	HGLLPCLHVA	ATTUTTEREM	-844
	HSV1-Patton	SSPEEAVLLD	MOON Y TEAM O	MOVIGE IGV	HGLLPCLHVA	ATVITIONEM	-844
	HSV1-DJL	SSPEEAVLLD	KOOAATKVVC	MOVIGETGV	HGLLPCLHVA	Antimatcrem	-844
55	HSV1-F	SSPEEAVLLD	KÕÕHHTKAAC	MSVIGFIGVQ	, ugnilectiva	AIVIIIGICA	044
	HSV2-MS	LLATRAYVHA	RWAEFDQLLA	DFPEAAGMRA	PGPYSMRIIY	GDTDSIFVLC	-897
	HSV2-186	LLATRAYVHA	RWAEFDOLLA	DFPEAAGMRA	PGPYSMRIIY	GDTDSIFVLC	-899
	HSV-Kos	LLATREYVHA	RWAAFEOLLA	DFPEAADMRA	PGPYSMRIIY	GDTDSIFVLC	-894
60	HSV1-Patton	LLATREYVHA	RWAAFEOLLA	DFPEAADMRA	PGPYSMRIIY	GDTDSIFVLC	-894
UU	HSV1-DJL	LLATREYVHA	RWAAFEOLLA	DFPEAADMRA	A PGPYSMRIIY	GDTDSIFVLC	-894
	HSV1-F	LLATREYVHA	RWAAFEQLLA	DFPEAADMRA	PGPYSMRIIY	GDTDSIFVLC	-894
	HSV2-MS	RGLTAAGLVA	MGDKMASHIS	RALFLPPIKI	ECEKTFTKLL	LIAKKKYIGV	_010
65	HSV2-186	RGLTAAGLVA	MGUKMASHIS	KALFLPPIKI	ECEKTFTKLL	LTAKKKXIGV	_949 _0//
	HSV-Kos	RGLTAAGLTA	MGUKMASHIS	KALFLPPIKI	ECEKTFTKLL	T TAKKKY TGV	_0// -344
	HSV1-Patton	RGLTAAGLTA	MGUKMASHIS	KALFLPPIKI	ECEKTFTKLL	PIAKKKXIGA	-344

	HSV1-DJL	RGLTAAGLTA	ACDKWY CHIC	RALFLPPIKL	ECEKTFTKLL	LIAKKKYIGV	-944
	HSV1-F	RGLTAAGLTA		, <u>, , , , , , , , , , , , , , , , , , </u>		LIAKKKYIGV	-944
	U2AT I	KODIAKOBIA	V CDIGHIDING				
	HSV2-MS	ICGGKMLIKG	VDLVRKNNCA	FINRTSRALV	DLLFYDDTVS	GAAAALAERP	- 997
5	HSV2-186		VDLVRKNNCA				-999
,	HSV-Kos	IYGGKMLIKG	VDLVRKNNCA	FINRTSRALV	DLLFYDDTVS	GAAAALAERP	-994
	HSV1-Patton	IYGGKMLIKG	VDLVRKNNCA	FINRTSRALV	DLLFYDDTVS	GAAAALAERP	-994
	HSV1-DJL	IYGGKMLIKG	VDLVRKNNCA	FINRTSRALV	DLLFYDDTVS	GAAAALAERP	-994
	HSV1-F	IYGGKMLIKG	VDLVRKNNCA	FINRTSRALV	DLLFYDDTVS	GAAAALAERP	-994
10							
	HSV2-MS	AEEWLARPLP	EGLQAFGAVL	VDAHRRITDP	ERDIQDFVLT	AELSRHPRAY	-1047
	HSV2-186	AEEWLARPLP	EGLQAFGAVL		ERDIQDFVLT	AELSRHPRAY	-1049
	HSV-Kos	AEEWLARPLP	EGLQAFGAVL		ERDIQDFVLT	AELSRHPRAY	-1044
	HSV1-Patton	AEEWLARPLP	EGLQAFGAVL		ERDIQDFVLT	AELSRHPRAY	-1044
15	HSV1-DJL	AEEWLARPLP	EGLQAFGAVL		ERDIQDFVLT	AELSRHPRAY	-1044
10	HSV1-F	AEEWLARPLP	EGLQAFGAVL		ERDIQDFVLT	AELSRHPRAY	-1044
	11012 2		2				
	HSV2-MS	TNKRLAHLTV	YYKLMARRAQ	VPSIKDRIPY	VIVAQTREVE	ETVARLAALR	-1097
	HSV2-186		YYKLMARRAQ			ETVARLAALR	-1099
20	HSV-Kos		YYKLMARRAQ			ETVARLAALR	-1094
	HSV1-Patton		YYKLMARRAQ			ETVARLAALR	-1094
	HSV1-DJL	TNKRLAHLTV	YYKLMARRAQ	VPSIKDRIPY	VIVAQTREVE	ETVARLAALR	-1094
	HSV1-F		YYKLMARRAQ			ETVARLAALR	-1094
25	HSV2-MS	ELDAAAPGDE	PAPPAALPSP	AKRPRETPSH	ADPPGGASKP	RKLLVSELAE	-1147
	HSV2-186	ELDAAAPGDE	PAPPAALPSP	AKRPRETPSH	ADPPGGASKP	RKLLVSELAE	-1149
	HSV-Kos				ADPPGGASKP		
	HSV1-Patton				ADPPGGASKP		
	HSV1-DJL				ADPPGGASKP		
30	HSV1-F	ELDAAAPGDE	PAPPAALPSP	AKRPRETPLH	ADPPGGASKP	RKLLVSELAE	-1144
	HSV2-MS	DPGYAIARGV	PLNTDYYFSH	LLGAACVTFK	ALFGNNAKIT	ESLLKRFIPE	-1197
	HSV2-186	DPGYAIARGV	PLNTDYYFSH	LLGAACVTFK	ALFGNNAKIT	ESLLKRFIPE	-1199
	HSV-Kos				ALFGNNAKIT		
35	HSV1-Patton				ALFGNNAKIT		
	HSV1-DJL				ALFGNNAKIT		
	HSV1-F	DPAYAIAHGV	ALNTDYYFSH	LLGAACVTFK	ALFGNNAKIT	ESLLKRFIPE	-1194
	HSV2-MS				RMLHRAFDTL		
40	HSV2-186				RMLHRAFDTL		
	HSV-Kos				RMLHRAFDTL		
	HSV1-Patton	VWHPPDDVTA	RLRAAGFGAV	GAGATAEETR	RMLHRAFDTL	A* -1235	
	HSV1-DJL				RMLHRAFDTL		
	HSV1-F	VWHPPDDVAA	RLRAAGFGAV	GAGATAEETR	RMLHRAFDTL	A* -1235	
45							
7.5							

^{*}Amino acid alignment demonstrates difference in amino acid's sequences.

^{*}The gaps "...." indicate missing amino acids relative to other stanins.

^{*}Wild HSV2-MS is listed as SEQ. ID NO 14.

^{*}Wild HSV2-186 is listed as SEQ. ID NO 15.

^{*}Wild HSV-Kos is listed as SEQ. ID NO 16.

^{*}Wild HSV1-Patton is listed as SEQ. ID NO 17.

^{*}Wild HSV1-DJL is listed as SEQ. ID NO 18.

^{*}Wild HSV1-F is listed as SEQ. ID NO 19.

Figure 5 DNA and amino acid sequence list

SEQ. ID. NO. 1 DNA sequence of DNA polymerase gene for HSV2-MS-M1

1 ATGTTTTGTG CCGCGGGCGG CCCGACTTCC CCCGGGGGGA AGTCGGCGGC 5 51 TCGGGCGGCG TCTGGGTTTT TTGCCCCCCA CAACCCCCGG GGAGCCACCC 101 AGACGGCACC GCCGCCTTGC CGCCGGCAGA ACTTCTACAA CCCCCACCTC 10 151 GCTCAGACCG GAACGCAGCC AAAGGCCCCC GGGCCGGCTC AGCGCCATAC 201 GTACTACAGC GAGTGCGACG AATTTCGATT TATCGCCCCG CGTTCGCTGG 251 ACGAGGACGC CCCCGCGGAG CAGCGCACCG GGGTCCACGA CGGCCGCCTC 15 301 CGGCGCCC CTAAGGTGTA CTGCGGGGGG GACGAGCGCG ACGTCCTCCG 351 CGTGGGCCCG GAGGGCTTCT GGCCGCGTCG CTTGCGCCCTG TGGGGCGGTG 20 401 CGGACCATGC CCCCAAGGGG TTCGACCCCA CCGTCACCGT CTTCCACGTG 451 TACGACATCC TGGAGCACGT GGAACACGCG TACAGCATGC GCGCCGCCCA 501 GCTCCACGAG CGATTTATGG ACGCCATCAC GCCCGCCGGG ACCGTCATCA 25 551 CGCTTCTGGG TCTGACCCCC GAAGGCCATC GCGTCGCCGT TCACGTCTAC 601 GGCACGCGC AGTACTTTTA CATGAACAAG GCGGAGGTGG ATCGGCACCT 30 651 GCAGTGCCGT GCCCCGCGCG ATCTCTGCGA GCGCCTGGCG GCGGCCCTGC 701 GCGAGTCGCC GGGGGCGTCG TTCCGCGGCA TCTCCGCGGA CCACTTCGAG 751 GCGGAGGTGG TGGAGCGCGC CGACGTGTAC TATTACGAAA CGCGCCCGAC 35 801 CCTGTACTAC CGCGTCTTCG TGCGAAGCGG GCGCGCGCTG GCCTACCTGT 851 GCGACAACTT TTGCCCCGCG ATCAGGAAGT ACGAGGGGGG CGTCGACGCC 40 901 ACCACCGGT TTATCCTGGA CAACCCGGGG TTTGTCACCT TCGGCTGGTA 951 CCGCCTCAAG CCCGGCCGCG GGAACGCGCC GGCCCAACCG CGCCCCCGA 1001 CGGCGTTCGG AACCTCGAGC GACGTCGAGT TTAACTGCAC GGCGGACAAC 45 1051 CTGGCCGTCG AGGGGGCCAT GTGTGACCTG CCGGCCTACA AGCTCATGTG 1101 CTTCGATATC GAATGCAAGG CCGGGGGGGA GGACGAGCTG GCCTTTCCGG 50 1151 TCGCGGAACG CCCGGAAGAC CTCGTCATCC AGATCTCCTG TCTGCTCTAC 1201 GACCTGTCCA CCACCGCCCT CGAGCACATC CTCCTGTTTT CGCTCGGATC 1251 CTGCGACCTC CCCGAGTCCC ACCTCAGCGA TCTCGCCTCC AGGGGCCTGC 55 1301 CGGCCCCCGT CGTCCTGGAG TTTGACAGCG AATTCGAGAT GCTGCTGGCC

1351 TTCATGACCT TCGTCAAGCA GTACGGCCCC GAGTTCGTGA CCGGGTACAA 1401 CATCATCAAC TTCGACTGGC CCTTCGTCCT GACCAAGCTG ACGGAGATCT 1451 ACAAGGTCCC GCTCGACGGG TACGGGCGCA TGAACGGCCG GGGTGTGTTC 5 1501 CGCGTGTGGG ACATCGGCCA GAGCCACTTT CAGAAGCGCA GCAAGATCAA 1551 GGTGAACGGG ATGGTGAACA TCGACATGTA CGGCATCATC ACCGACAAGG 10 1601 TCAAACTCTC CAGCTACAAG CTGAACGCCG TCGCCGAGGC CGTCTTGAAG 1651 GACAAGAAGA AGGATCTGAG CTACCGCGAC ATCCCCGCCT ACTACGCCTC 1701 CGGGCCCGCG CAGCGCGGGG TGATCGGCGA GTATTGTGTG CAGGACTCGC 15 1751 TGCTGGTCGG GCAGCTGTTC TTCAAGTTTC TGCCGCACCT GGAGCTTTCC 1801 GCCGTCGCGC GCCTGGCGGG CATCAACATC ACCCGCACCA TCTACGACGG 20 1851 CCAGCAGATC CGCGTCTTCA CGTGCCTCCT GCGCCTTGCG GGCCAGAAGG 1901 GCTTCATCCT GCCGGACACC CAGGGGCGGT TTCGGGGCCT CGACAAGGAG 1951 GCGCCCAAGC GCCCGGCCGT GCCTCGGGGG GAAGGGGAGC GGCCGGGGGA 25 2001 CGGGAACGGG GACGAGGATA AGGACGACGA CGAGGACGAG GACGGGGACG 2051 AGCGCGAGGA GGTCGCGCGC GAGACCGGGG GCCGGCACGT TGGGTACCAG 30 2101 GGGGCCCGGG TCCTCGACCC CACCTCCGGG TTTCACGTCG ACCCCGTGGT 2151 GGTGTTTGAC TTTGCCAGCC TGTACCCCAG CATCATCCAG GCCCACAACC 2201 TGTGCTTCAG TACGCTCTCC CTGCGGCCCG AGGCCGTCGC GCACCTGGAG 35 2251 GCGGACCGGG ACTACCTGGA GATCGAGGTG GGGGGCCGAC GGCTGTTCTT 2301 CGTGAAGGCC CACGTACGCG AGAGCCTGCT GAGCATCCTG CTGCGCGACT 40 2351 GGCTGGCCAT GCGAAAGCAG ATCCGCTCGC GGATCCCCCA GAGCACCCCC 2401 GAGGAGGCCG TCCTCCTCGA CAAGCAACAG GCCGCCATCA AGGTGGTGTG 45 2501 TGCACGTGGC CGCCACCGTG ACGACCATCG GCCGCGAGAT GCTCCTCGCG 2551 ACGCGCGCGT ACGTGCACGC GCGCTGGGCG GAGTTCGATC AGCTGCTGGC 50 2601 CGACTTTCCG GAGGCGGCCG GCATGCGCGC CCCCGGTCCG TACTCCATGC 2651 GCATCATCTA CGGGGACACG GACTCCATTT TCGTTTTGTG CCGCGGCCTC 2701 ACGCCCGCGG GCCTGGTGGC CATGGGCGAC AAGATGGCGA GCCACATCTC 55 2751 GCGCGCGCTG TTCCTCCCCC CGATCAAGCT CGAGTGCGAA AAAACGTTCA 2801 CCAAGCTGCT GCTCATCGCC AAGAAAAAGT ACATCGGCGT CATCTGCGGG 60

	851 GGCAAGATGC TCATCAAGGG CGTGGATCTG GTGCGCAAAA ACAACTGCGC
	901 GTTTATCAAC CGCACCTCCA GGGCCCTGGT CGACCTGCTG TTTTACGACG
5	951 ATACCGTATC CGGAGCGCC GCCGCGTTAG CCGAGCGCCC CGCAGAGGAC
	1001 TGGCTGGCGC GACCCCTGCC CGAGGGACTG CAGGCGTTCG GGGCCGTCCT
10	051 CGTAGACGCC CATCGGCGCA TCACCGACCC GGAGAGGGAC ATCCAGGACT
10	101 TTGTCCTCAC CGCCGAACTG AGCAGACACC CGCGCGCGTA CACCAACAAG
	151 CGCCTGGCCC ACCTGACGGT GTATTACAAG CTCATGGCCC GCCGCGCGCA
15	3201 GGTCCCGTCC ATCAAGGACC GGATCCCGTA CGTGATCGTG GCCCAGACCC
	3251 GCGAGGTAGA GGAGACGGTC GCGCGGCTGG CCGCCCTCCG CGAGCTAGA
20	3301 GCCGCCGCCC CAGGGGACGA GCCCGCCCCC CCAGCGGCCC TGCCCTCCCC
20	351 GGCCAAGCGC CCCCGGGAGA CGCCGTCGCA TGCCGACCCC CCGGGAGGCC
	3401 CGTCCAAGCC CCGCAAGCTG CTGGTGTCCG AGCTGGCGGA GGATCCCGGC
25	3451 TACGCCATCG CCCGGGGCGT TCCGCTCAAC ACGGACTATT ACTTCTCGCA
	3501 CCTGCTGGGG GCGGCCTGCG TGACGTTCAA GGCCCTGTTT GGAAATAACC
30	3551 CCAAGATCAC CGAGAGTCTG TTAAAGAGGT TTATTCCCGA GACGTGGCAC
50	3601 CCCCCGGACG ACGTGGCCGC GCGGCTCAGG GCCGCGGGGT TCGGGCCGG
	3651 GGGGCCGGC GCTACGGCGG AGGAAACTCG TCGAATGTTG CATAGAGCC
35	3701 TTGATACTCT AGCATGA

SEQ. ID. NO. 2 Amino acid sequence of DNA polymerase for HSV2-MS-M1

	1 MFCAAGGPTS PGGKSAARAA SGFFAPHNPR GATQTAPPPC RRQNFYNPHL
5	51 AQTGTQPKAP GPAQRHTYYS ECDEFRFIAP RSLDEDAPAE QRTGVHDGRL
	101 RRAPKVYCGG DERDVLRVGP EGFWPRRLRL WGGADHAPKG FDPTVTVFHV
10	151 YDILEHVEHA YSMRAAQLHE RFMDAITPAG TVITLLGLTP EGHRVAVHVY
	201 GTRQYFYMNK AEVDRHLQCR APRDLCERLA AALRESPGAS FRGISADHFE
	251 AEVVERADVY YYETRPTLYY RVFVRSGRAL AYLCDNFCPA IRKYEGGVDA
15	301 TTRFILDNPG FVTFGWYRLK PGRGNAPAQP RPPTAFGTSS DVEFNCTADN
	351 LAVEGAMCDL PAYKLMCFDI ECKAGGEDEL AFPVAERPED LVIQISCLLY
20	401 DLSTTALEHI LLFSLGSCDL PESHLSDLAS RGLPAPVVLE FDSEFEMLLA
20	451 FMTFVKQYGP EFVTGYNIIN FDWPFVLTKL TEIYKVPLDG YGRMNGRGVF
	501 RVWDIGQSHF.QKRSKIKVNG MVNIDMYGII TDKVKLSSYK LNAVAEAVLK
25	551 DKKKDLSYRD IPAYYASGPA QRGVIGEYCV QDSLLVGQLF FKFLPHLELS
	601 AVARLAGINI TRTIYDGQQI RVFTCLLRLA GQKGFILPDT QGRFRGLDKE
30	651 APKRPAVPRG EGERPGDGNG DEDKDDDEDE DGDEREEVAR ETGGRHVGYC
50	701 GARVLDPTSG FHVDPVVVFD FASLYPSIIQ AHNLCFSTLS LRPEAVAHLE
	751 ADRDYLEIEV GGRRLFFVKA HVRESLLSIL LRDWLAMRKQ IRSRIPQSTP
35	801 EEAVLLDKQQ AAIKVVCNSV YGFTGAQHGL LPCLHVAATV TTIGREMLLA
	851 TRAYVHARWA EFDQLLADFP EAAGMRAPGP YSMRIIYGDT DSIFVLCRGL
40	901 TAAGLVAMGD KMASHISRAL FLPPIKLECE KTFTKLLLIA KKKYIGVICG
.0	951 GKMLIKGVDL VRKNNCAFIN RTSRALVDLL FYDDTVSGAA AALAERPAEE
	1001 WLARPLPEGL QAFGAVLVDA HRRITDPERD IQDFVLTAEL SRHPRAYTNK
45	1051 RLAHLTVYYK LMARRAQVPS IKDRIPYVIV AQTREVEETV ARLAALRELD
	1101 AAAPGDEPAP PAALPSPAKR PRETPSHADP PGGASKPRKL LVSELAEDPG
50	1151 YAIARGVPLN TDYYFSHLLG AACVTFKALF GNNAKITESL LKRFIPETWH
20	1201 PPDDVAARLR AAGFGPAGAG ATAEETRRML HRAFDTLA*

DNA sequence of DNA polymerase gene for HSV2-186-M1 SEQ.ID.NO. 3 1 ATGTTTTGTG CCGCGGCGG CCCGGCTTCC CCCGGGGGA AGTCGGCGGC 51 TCGGGCGCG TCTGGGTTTT TTGCCCCCCA CAACCCCCGG GGAGCCACCC 5 101 AGACGGCACC GCCGCCTTGC CGCCGGCAGA ACTTCTACAA CCCCCACCTC 151 GCTCAGACCG GAACGCAGCC AAAGGCCCCC GGGCCGGCTC AGCGCCATAC 10 201 GTACTACAGC GAGTGCGACG AATTTCGATT TATCGCCCCG CGTTCGCTGG 251 ACGAGGACGC CCCCGCGGAG CAGCGCACCG GGGTCCACGA CGGCCGCCTC 301 CGGCGCCCC CTAAGGTGTA CTGCGGGGGG GACGAGCGCG ACGTCCTCCG 15 351 CGTGGGCCCG GAGGGCTTCT GGCCGCGTCG CTTGCGCCTG TGGGGCGGTG 401 CGGACCATGC CCCCGAGGGG TTCGACCCCA CCGTCACCGT CTTCCACGTG 20 451 TACGACATCC TGGAGCACGT GGAACACGCG TACAGCATGC GCGCCGCCCA 501 GCTCCACGAG CGATTTATGG ACGCCATCAC GCCCGCCGGG ACCGTCATCA 551 CGCTTCTGGG TCTGACCCCC GAAGGCCATC GCGTCGCCGT TCACGTCTAC 25 601 GGCACGCGC AGTACTTTTA CATGAACAAG GCGGAGGTGG ATCGGCACCT 651 GCAGTGCCGT GCCCCGCGC ATCTCTGCGA GCGCCTGGCG GCGGCCCTGC 30 701 GCGAGTCGCC GGGGGCGTCG TTCCGCGGCA TCTCCGCGGA CCACTTCGAG 751 GCGGAGGTGG TGGAGCGCGC CGACGTGTAC TATTACGAAA CGCGCCCGAC 801 CCTGTACTAC CGCGTCTTCG TGCGAAGCGG GCGCGCGCTG GCCTACCTGT 35 851 GCGACAACTT TTGCCCCGCG ATCAGGAAGT ACGAGGGGGG CGTCGACGCC 901 ACCACCGGT TTATCCTGGA CAACCGGGG TTTGTCACCT TCGGCTGGTA 40 951 CCGCCTCAAG CCCGGCCGCG GGAACGCGCC GGCCCAACCG CGCCCCCGA 1001 CGGCGTTCGG AACCTCGAGC GACGTCGAGT TTAACTGCAC GGCGGACAAC 1051 CTGGCCGTCG AGGGGGCCAT GTGTGACCTG CCGGCCTACA AGCTCATGTG 45 1101 CTTCGATATC GAATGCAAGG CCGGGGGGGA GGACGAGCTG GCCTTTCCGG 1151 TCGCGGAACG CCCGGAAGAC CTCGTCATCC AGATCTCCTG TCTGCTCTAC 50 1201 GACCTGTCCA CCACCGCCCT CGAGCACATC CTCCTGTTTT CGCTCGGATC 1251 CTGCGACCTC CCCGAGTCCC ACCTCAGCGA TCTCGCCTCC AGGGGCCTGC 1301 CGGCCCCCGT CGTCCTGGAG TTTGACAGCG AATTCGAGAT GCTGCTGGCC 55 1351 TTCATGACCT TCGTCAAGCA GTACGGCCCC GAGTTCGTGA CCGGGTACAA 1401 CATCATCAAC TTCGACTGGC CCTTCGTCCT GACCAAGCTG ACGGAGATCT 60

	1451	ACAAGGTCCC GCTCGACGGG TACGGGCGCA TGAACGGCCG GGGTGTGTTC
	1501	CGCGTGTGGG ACATCGGCCA GAGCCACTTT CAGAAGCGCA GCAAGATCAA
5	1551	GGTGAACGGG ATGGTGAACA TCGACATGTA CGGCATCATC ACCGACAAGG
	1601	TCAAACTCTC CAGCTACAAG CTGAACGCCG TCGCCGAGGC CGTCTTGAAG
10	1651	GACAAGAAGA AGGATCTGAG CTACCGCGAC ATCCCCGCCT ACTACGCCTC
10	1701	CGGGCCCGCG CAGCGCGGG TGATCGGCGA GTATTGTGTG CAGGACTCGC
	1751	TGCTGGTCGG GCAGCTGTTC TTCAAGTTTC TGCCGCACCT GGAGCTTTCC
15	1801	GCCGTCGCGC GCCTGGCGGG CATCAACATC ACCCGCACCA TCTACGACGG
	1851	CCAGCAGATC CGCGTCTTCA CGTGCCTCCT GCGCCTTGCG GGCCAGAAGG
20	1901	GCTTCATCCT GCCGGACACC CAGGGGCGGT TTCGGGGCCT CGACAAGGAG
	1951	GCGCCCAAGC GCCCGGCCGT GCCTCGGGGG GAAGGGGAGC GGCCGGGGA
	2001	CGGGAACGGG GACGAGGATA AGGACGACGA CGAGGACGG GACGAGGACG
25	2051	GGGACGAGCG CGAGGAGGTC GCGCGCGAGA CCGGGGGCCG GCACGTTGGG
	2101	TACCAGGGG CCCGGGTCCT CGACCCCACC TCCGGGTTTC ACGTCGACCC
30	2151	CGTGGTGGTG TTTGACTTTG CCAGCCTGTA CCCCAGCATC ATCCAGGCCC
30	2201	ACAACCTGTG CTTCAGTACG CTCTCCCTGC GGCCCGAGGC CGTCGCGCAC
	2251	CTGGAGGCGG ACCGGGACTA CCTGGAGATC GAGGTGGGGG GCCGACGGCT
35	2301	GTTCTTCGTG AAGGCCCACG TACGCGAGAG CCTGCTGAGC ATCCTGCTGC
	2351	GCGACTGGCT GGCCATGCGA AAGCAGATCC GCTCGCGGAT CCCCCAGAGC
40	2401	CCCCCGAGG AGGCCGTCCT CCTCGACAAG CAACAGGCCG CCATCAAGGT
40	2451	GGTGTGCAAC TCGGTGTACG GGTTCACCGG GGCGCAGCAC GGTCTTCTGC
	2501	CCTGCCTGCA CGTGGCCGC ACCGTGACGA CCATCGGCCG CGAGATGCTC
45	2551	CTCGCGACGC GCGCGTACGT GCACGCGCGC TGGGCGGAGT TCGATCAGCT
	2601	GCTGGCCGAC TTTCCGGAGG CGGCCGGCAT GCGCGCCCCC GGTCCGTACT
50	2651	CCATGCGCAT CATCTACGGG GACACGGACT CCATTTTCGT TTTGTGCCGC
30	2701	GGCCTCACGG CCGCGGGCCT GGTGGCCATG GGCGACAAGA TGGCGAGCCA
	2751	CATCTCGCGC GCGCTGTTCC TCCCCCCGAT CAAGCTCGAG TGCGAAAAAA
55	2801	CGTTCACCAA GCTGCTGCTC ATCGCCAAGA AAAAGTACAT CGGCGTCATC
	2851	TGCGGGGCA AGATGCTCAT CAAGGGCGTG GATCTGGTGC GCAAAAACAA
60	2901	CTGCGCGTTT ATCAACCGCA CCTCCAGGGC CCTGGTCGAC CTGCTGTTTT
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2951 ACGACGATAC CGTATCCGGA GCGCCCGCG CGTTAGCCGA GCGCCCCGCA 3001 GAGGAGTGGC TGGCGCGACC CCTGCCCGAG GGACTGCAGG CGTTCGGGGC 3051 CGTCCTCGTA GACGCCCATC GGCGCATCAC CGACCCGGAG AGGGACATCC 3101 AGGACTTTGT CCTCACCGCC GAACTGAGCA GACACCCGCG CGCGTACACC 3151 AACAAGCGCC TGGCCCACCT GACGGTGTAT TACAAGCTCA TGGCCCGCCG 3201 CGCGCAGGTC CCGTCCATCA AGGACCGGAT CCCGTACGTG ATCGTGGCCC 3251 AGACCCGCGA GGTAGAGGAG ACGGTCGCGC GGCTGGCCGC CCTCCGCGAG 3301 CTAGACGCCG CCGCCCCAGG GGACGAGCCC GCCCCCCAG CGGCCCTGCC 3351 CTCCCCGGCC AAGCGCCCCC GGGAGACGCC GTCGCATGCC GACCCCCCGG 3401 GAGGCGCGTC CAAGCCCCGC AAGCTGCTGG TGTCCGAGCT GGCGGAGGAT 3451 CCCGGGTACG CCATCGCCCG GGGCGTTCCG CTCAACACGG ACTATTACTT 3501 CTCGCACCTG CTGGGGGCGG CCTGCGTGAC GTTCAAGGCC CTGTTTGGAA 3551 ATAACGCCAA GATCACCGAG AGTCTGTTAA AGAGGTTTAT TCCCGAGACG 3601 TGGCACCCCC CGGACGACGT GGCCGCGCGC CTCAGGGCCG CGGGGTTCGG 3651 GCCGGCGGGGGCGCTA CGGCGGAGGA AACTCGTCGA ATGTTGCATA 3701 GAGCCTTTGA TACTCTAGCA TGA

Amino acid sequence of DNA polymerase for HSV2-186-M1 SEQ.ID.NO. 4 1 MFCAAGGPAS PGGKSAARAA SGFFAPHNPR GATQTAPPPC RRQNFYNPHL 5 51 AQTGTQPKAP GPAQRHTYYS ECDEFRFIAP RSLDEDAPAE QRTGVHDGRL 101 RRAPKVYCGG DERDVLRVGP EGFWPRRLRL WGGADHAPEG FDPTVTVFHV 10 151 YDILEHVEHA YSMRAAQLHE RFMDAITPAG TVITLLGLTP EGHRVAVHVY 201 GTRQYFYMNK AEVDRHLQCR APRDLCERLA AALRESPGAS FRGISADHFE 251 AEVVERADVY YYETRPTLYY RVFVRSGRAL AYLCDNFCPA IRKYEGGVDA 15 301 TTRFILDNPG FVTFGWYRLK PGRGNAPAQP RPPTAFGTSS DVEFNCTADN 351 LAVEGAMCDL PAYKLMCFDI ECKAGGEDEL AFPVAERPED LVIQISCLLY 20 401 DLSTTALEHI LLFSLGSCDL PESHLSDLAS RGLPAPVVLE FDSEFEMLLA 451 FMTFVKQYGP EFVTGYNIIN FDWPFVLTKL TEIYKVPLDG YGRMNGRGVF 501 RVWDIGQSHF QKRSKIKVNG MVNIDMYGII TDKVKLSSYK LNAVAEAVLK 25 551 DKKKDLSYRD IPAYYASGPA QRGVIGEYCV QDSLLVGQLF FKFLPHLELS 601 AVARLAGINI TRTIYDGQQI RVFTCLLRLA GQKGFILPDT QGRFRGLDKE 30 651 APKRPAVPRG EGERPGDGNG DEDKDDDEDG DEDGDEREEV ARETGGRHVG 701 YQGARVLDPT SGFHVDPVVV FDFASLYPSI IQAHNLCFST LSLRPEAVAH 751 LEADRDYLEI EVGGRRLFFV KAHVRESLLS ILLRDWLAMR KQIRSRIPQS 35 801 PPEEAVLLDK QQAAIKVVCN SVYGFTGAQH GLLPCLHVAA TVTTIGREML 851 LATRAYVHAR WAEFDQLLAD FPEAAGMRAP GPYSMRIIYG DTDSIFVLCR 40 901 GLTAAGLVAM GDKMASHISR ALFLPPIKLE CEKTFTKLLL IAKKKYIGVI 951 CGGKMLIKGV DLVRKNNCAF INRTSRALVD LLFYDDTVSG AAAALAERPA 1001 EEWLARPLPE GLQAFGAVLV DAHRRITDPE RDIQDFVLTA ELSRHPRAYT 45 1051 NKRLAHLTVY YKLMARRAQV PSIKDRIPYV IVAQTREVEE TVARLAALRE 1101 LDAAAPGDEP APPAALPSPA KRPRETPSHA DPPGGASKPR KLLVSELAED 50 1151 PGYAIARGVP LNTDYYFSHL LGAACVTFKA LFGNNAKITE SLLKRFIPET 1201 WHPPDDVAAR LRAAGFGPAG AGATAEETRR MLHRAFDTLA *

DNA sequence of DNA polymerase gene for HSV1-KOS-M1 SEQ.ID.NO. 5 1 ATGTTTTCCG GTGGCGGCGG CCCGCTGTCC CCCGGAGGAA AGTCGGCGGC 51 CAGGGCGCG TCCGGGTTTT TTGCGCCCGC CGGCCCTCGC GGAGCCGGCC 5 101 GGGGACCCCC GCCTTGTTTG AGGCAAAACT TTTACAACCC CTACCTCGCC 151 CCAGTCGGGA CGCAACAGAA GCCGACCGGG CCAACCCAGC GCCATACGTA 10 201 CTATAGCGAA TGCGATGAAT TTCGATTCAT CGCCCCGCGG GTGCTGGACG 251 AGGATGCCCC CCCGGAGAAG CGCGCCGGGG TGCACGACGG TCACCTCAAG 301 CGCGCCCCA AGGTGTACTG CGGGGGGGAC GAGCGCGACG TCCTCCGCGT 15 351 CGGGTCGGGC GGCTTCTGGC CGCGGCGCTC GCGCCTGTGG GGCGGCGTGG 401 ACCACGCCCC GGCGGGGTTC AACCCCACCG TCACCGTCTT TCACGTGTAC 20 451 GACATCCTGG AGAACGTGGA GCACGCGTAC GGCATGCGCG CGGCCCAGTT 501 CCACGCGCGG TTTATGGACG CCATCACACC GACGGGGACC GTCATCACGC 551 TCCTGGGCCT GACTCCGGAA GGCCACCGGG TGGCCGTTCA CGTTTACGGC 25 601 ACGCGGCAGT ACTTTTACAT GAACAAGGAG GAGGTTGACA GGCACCTACA 651 ATGCCGCGCC CCACGAGATC TCTGCGAGCG CATGGCCGCG GCCCTGCGCG 30 701 AGTCCCCGGG CGCGTCGTTC CGCGGCATCT CCGCGGACCA CTTCGAGGCG 751 GAGGTGGTGG AGCGCACCGA CGTGTACTAC TACGAGACGC GCCCCGCTCT 801 GTTTTACCGC GTCTACGTCC GAAGCGGGCG CGTGCTGTCG TACCTGTGCG 35 851 ACAACTTCTG CCCGGCCATC AAGAAGTACG AGGGTGGGGT CGACGCCACC 901 ACCCGGTTCA TCCTGGACAA CCCCGGGTTC GTCACCTTCG GCTGGTACCG 40 951 TCTCAAACCG GGCCGGAACA ACACGCTAGC CCAGCCGCGG GCCCCGATGG 1001 CCTTCGGGAC ATCCAGCGAC GTCGAGTTTA ACTGTACGGC GGACAACCTG 1051 GCCATCGAGG GGGGCATGAG CGACCTACCG GCATACAAGC TCATGTGCTT 45 1101 CGATATCGAA TGCAAGGCGG GGGGGGAGGA CGAGCTGGCC TTTCCGGTGG 1151 CCGGGCACCC GGAGGACCTG GTTATTCAGA TATCCTGTCT GCTCTACGAC 50 1201 CTGTCCACCA CCGCCCTGGA GCACGTCCTC CTGTTTTCGC TCGGTTCCTG 1251 CGACCTCCCC GAATCCCACC TGAACGAGCT GGCGGCCAGG GGCCTGCCCA 1301 CGCCCGTGGT TCTGGAATTC GACAGCGAAT TCGAGATGCT GTTGGCCTTC 55 1351 ATGACCCTTG TGAAACAGTA CGGCCCCGAG TTCGTGACCG GGTACAACAT 1401 CATCAACTTC GACTGGCCCT TCTTGCTGGC CAAGTTGACG GACATTTACA

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1451 AGGTCCCCCT GGACGGGTAC GGCCGCATGA ACGGCCGGGG CGTGTTTCGC 1501 GTGTGGGACA TAGGCCAGAG CCACTTCCAG AAGCGCAGCA AGATAAAGGT 1551 GAACGCCATG GTGAACATCG ACATGTACGG GATCATAACC GACAAGATCA 1601 AGCTCTCGAG CTACAAGCTC AACGCCGTGG CCGAAGCCGT CCTGAAGGAC 1651 AAGAAGAAGG ACCTGAGCTA TCGCGACATC CCCGCCTACT ACGCCGCCGG 1701 GCCCGCGCAA CGCGGGGTGA TCGGCGAGTA CTGCATACAG GATTCCCTGC 1751 TGGTGGCCA GCTGTTTTTT AAGTTTTTGC CCCATCTGGA GCTCTCGGCC 1801 GTCGCGCGCT TGGCGGGTAT TAACATCACC CGCACCATCT ACGACGCCA 1851 GCAGATCCGC GTCTTTACGT GCCTGCTGCG CCTGGCCGAC CAGAAGGGCT 1901 TTATTCTGCC GGACACCCAG GGGCGATTTA GGGGCGCCGG GGGGGAGGCG 1951 CCCAAGCGTC CGGCCGCAGC CCGGGAGGAC GAGGAGCGC CAGAGGAGGA 2001 GGGGGAGGAC GAGGACGAAC GCGAGGAGGG CGGGGGCGAG CGGGAGCCGG 2051 AGGGCGCGCGGGAGACCGCC GGCCGGCACG TGGGGTACCA GGGGGCCAGG 2101 GTCCTTGACC CCACTTCCGG GTTTCACGTG AACCCCGTGG TGGTGTTCGA 2151 CTTTGCCAGC CTGTACCCCA GCATCATCCA GGCCCACAAC CTGTGCTTCA 2201 GCACGCTCTC CCTGAGGGCC GACGCAGTGG CGCACCTGGA GGCGGCCAAG 2251 GACTACCTGG AGATCGAGGT GGGGGGGGGGGA CGGCTGTTCT TCGTCAAGGC 2301 TCACGTGCGA GAGAGCCTCC TCAGCATCCT CCTGCGGGAC TGGCTCGCCA 2351 TGCGAAAGCA GATCCGCTCG CGGATTCCCC AGAGCAGCCC CGAGGAGGCC 2401 GTGCTCCTGG ACAAGCAGCA GGCCGCCATC AAGGTCGTGT GTAACTCGGT 2451 GTACGGGTTC ACGGGAGCGC AGCACGGACT CCTGCCGTGC CTGCACGTTG 2501 CCGCGACGGT GACGACCATC GGCCGCGAGA TGCTGCTCGC GACCCGCGAG 2551 TACGTCCACG CGCGCTGGGC GGCCTTCGAA CAGCTCCTGG CCGATTTCCC 2601 GGAGGCGGCC GACATGCGCG CCCCCGGGCC CTATTCCATG CGCATCATCT 2651 ACGGGGACAC GGACTCCATA TTTGTGCTGT GCCGCGGCCT CACGGCCGCC 2701 GGGCTGACGG CCATGGGCGA CAAGATGGCG AGCCACATCT CGCGCGCGCT 2751 GTTTCTGCCC CCCATCAAAC TCGAGTGCGA AAAGACGTTC ACCAAGCTGC 2801 TGCTGATCGC CAAGAAAAG TACATCGGCG TCATCTACGG GGGTAAGATG 2851 CTCATCAAGG GCGTGGATCT GGTGCGCAAA AACAACTGCG CGTTTATCAA 2901 CCGCACCTCC AGGGCCCTGG TCGACCTGCT GTTTTACGAC GATACCGTAT

	2951 CCGGAGCGGC CGCCGCGTTA GCCGAGCGCC CCGCAGAGGA GTGGCTGGCG
	3001 CGACCCCTGC CCGAGGGACT GCAGGCGTTC GGGGCCGTCC TCGTAGACGC
5	3051 CCATCGGCGC ATCACCGACC CGGAGAGGGA CATCCAGGAC TTTGTCCTCA
	3101 CCGCCGAACT GAGCAGACAC CCGCGCGCGT ACACCAACAA GCGCCTGGCC
10	3151 CACCTGACGG TGTATTACAA GCTCATGGCC CGCCGCGCGC AGGTCCCGTC
10	3201 CATCAAGGAC CGGATCCCGT ACGTGATCGT GGCCCAGACC CGCGAGGTAG
	3251 AGGAGACGGT CGCGCGGCTG GCCGCCCTCC GCGAGCTAGA CGCCGCCGCC
15 .	3301 CCAGGGGACG AGCCCGCCCC CCCCGCGGCC CTGCCCTCCC CGGCCAAGCG
	3351 CCCCCGGGAG ACGCCGTCGC ATGCCGACCC CCCGGGAGGC GCGTCCAAGC
••	3401 CCCGCAAGCT GCTGGTGTCC GAGCTGGCCG AGGATCCCGC ATACGCCATT
20	3451 GCCCACGGCG TCGCCCTGAA CACGGACTAT TACTTCTCCC ACCTGTTGGG
	3501 GGCGGCGTGC GTGACATTCA AGGCCCTGTT TGGGAATAAC GCCAAGATCA
25	3551 CCGAGAGTCT GTTAAAAAGG TTTATTCCCG AAGTGTGGCA CCCCCCGGAC
	3601 GACGTGGCCG CGCGGCTCCG GGCCGCAGGG TTCGGGGCGG TGGGTGCCGC
20	3651 CGCTACGGCG GAGGAAACTC GTCGAATGTT GCATAGAGCC TTTGATACTC
30	3701 TAGCATGA

Amino acid sequence of DNA polymerase for HSV1-KOS-M1 SEQ.ID.NO. 6 1 MFSGGGGPLS PGGKSAARAA SGFFAPAGPR GAGRGPPPCL RQNFYNPYLA 5 51 PVGTQQKPTG PTQRHTYYSE CDEFRFIAPR VLDEDAPPEK RAGVHDGHLK 101 RAPKVYCGGD ERDVLRVGSG GFWPRRSRLW GGVDHAPAGF NPTVTVFHVY 151 DILENVEHAY GMRAAQFHAR FMDAITPTGT VITLLGLTPE GHRVAVHVYG 10 201 TRQYFYMNKE EVDRHLQCRA PRDLCERMAA ALRESPGASF RGISADHFEA 251 EVVERTDVYY YETRPALFYR VYVRSGRVLS YLCDNFCPAI KKYEGGVDAT 15 301 TRFILDNPGF VTFGWYRLKP GRNNTLAQPR APMAFGTSSD VEFNCTADNL 351 AIEGGMSDLP AYKLMCFDIE CKAGGEDELA FPVAGHPEDL VIQISCLLYD 401 LSTTALEHVL LFSLGSCDLP ESHLNELAAR GLPTPVVLEF DSEFEMLLAF 20 451 MTLVKQYGPE FVTGYNIINF DWPFLLAKLT DIYKVPLDGY GRMNGRGVFR 501 VWDIGQSHFQ KRSKIKVNGM VNIDMYGIIT DKIKLSSYKL NAVAEAVLKD 25 551 KKKDLSYRDI PAYYAAGPAQ RGVIGEYCIQ DSLLVGQLFF KFLPHLELSA 601 VARLAGINIT RTIYDGQQIR VFTCLLRLAD QKGFILPDTQ GRFRGAGGEA 651 PKRPAAARED EERPEEGED EDEREEGGGE REPEGARETA GRHVGYQGAR 30 701 VLDPTSGFHV NPVVVFDFAS LYPSIIQAHN LCFSTLSLRA DAVAHLEAGK 751 DYLEIEVGGR RLFFVKAHVR ESLLSILLRD WLAMRKQIRS RIPQSSPEEA 35 801 VLLDKQQAAI KVVCNSVYGF TGAQHGLLPC LHVAATVTTI GREMLLATRE 851 YVHARWAAFE QLLADFPEAA DMRAPGPYSM RIIYGDTDSI FVLCRGLTAA 901 GLTAMGDKMA SHISRALFLP PIKLECEKTF TKLLLIAKKK YIGVIYGGKM 40 951 LIKGVDLVRK NNCAFINRTS RALVDLLFYD DTVSGAAAAL AERPAEEWLA 1001 RPLPEGLQAF GAVLVDAHRR ITDPERDIQD FVLTAELSRH PRAYTNKRLA 45 1051 HLTVYYKLMA RRAQVPSIKD RIPYVIVAQT REVEETVARL AALRELDAAA 1101 PGDEPAPPAA LPSPAKRPRE TPSHADPPGG ASKPRKLLVS ELAEDPAYAI 1151 AHGVALNTDY YFSHLLGAAC VTFKALFGNN AKITESLLKR FIPEVWHPPD 50 1201 DVAARLRAAG FGAVGAGATA EETRRMLHRA FDTLA*

SEQ.ID.NO. 7 DNA sequence of HSV polymerase gene for HSV1-F-M1

5	1	ATGTTTTCCG	GTGGCGGCGG	CCCGCTGTCC	CCCGGAGGAA	AGTCGGCGGC
3	51	CAGGGCGGCG	TCCGGGTTTT	TTGCGCCCGC	CGGCCCTCGC	GGAGCCGGCC
	101	GGGGACCCCC	GCCTTGCTTG	AGGCAAAACT	TTTACAACCC	CTACCTCGCC
10	151	CCAGTCGGGA	CGCAACAGAA	GCCGACCGGG	CCAACCCAGC	GCCATACGTA
	201	CTATAGCGAA	TGCGATGAAT	TTCGATTCAT	CGCCCCGCGG	GTGCTGGACG
15	251	AGGATGCCCC	CCCGGAGAAG	CGCGCCGGGG	TGCACGACGG	TCACCTCAAG
12	301	CGCGCCCCCA	AGGTGTACTG	CGGGGGGGAC	GAGCGCGACG	TCCTCCGCGT
	351	CGGGTCGGGC	GGCTTCTGGC	CGCGGCGCTC	GCGCCTGTGG	GGCGGCGTGG
20	401	ACCACGCCCC	GGCGGGGTTC	AACCCCACCG	TCACCGTCTT	TCACGTGTAC
	451	GACATCCTGG	AGAACGTGGA	GCACGCGTAC	GGCATGCGCG	CGGCCCAGTT
25	501	CCACGCGCGG	TTTATGGACG	CCATCACACC	GACGGGGACC	GTCATCACGC
25	551	TCCTGGGCCT	GACTCCGGAA	GGCCACCGGG	TGGCCGTTCA	CGTTTACGGC
,	601	ACGCGGCAGT	ACTTTTACAT	GAACAAGGAG	GAGGTCGACA	GGCACCTACA
30	651	ATGCCGCGCC	CCACGAGATC	TCTGCGAGCG	CATGGCCGCG	GCCCTGCGCG
	701	AGTCCCCGGG	CGCGTCGTTC	CGCGGCATTT	CCGCGGACCA	CTTCGAGGCG
35	751	GAGGTGGTGG	AGCGCACCGA	CGTGTACTAC	TACGAGACGC	GCCCCGCTCT
55	801	GTTTTACCGC	GTCTACGTCC	GAAGCGGGCG	CGTGCTGTCG	TACCTGTGCG
	851	ACAACTTCTG	CCCGGCCATC	AAGAAGTACG	AGGGTGGGGT	CGACGCCACC
40	901	ACCCGGTTCA	TCCTGGACAA	CCCCGGGTTC	GTCACCTTCG	GCTGGTACCG
	951	TCTCAAACCG	GGCCGGAACA	ACACGCTAGC	CCAGCCGCGG	GCCCCGATGG
45	1001	CCTTCGGGAC	ATCCAGCGAC	GTCGAGTTTA	ACTGTACGGC	GGACAACCTG
43	1051	GCCATCGAGG	GGGGCATGAG	CGACCTACCG	GCATACAAGC	TCATGTGCTT
	1101	CGATATCGAA	TGCAAGGCGG	GGGGGGAGGA	CGAGCTGGCC	TTTCCGGTGG
50	1151	CCGGGCACCC	GGAGGACCTG	GTCATCCAGA	TATCCTGTCT	GCTCTACGAC
	1201	CTGTCCACCA	CCGCCCTGGA	GCACGTCCTC	CTGTTTTCGC	CTCGGTTCCTG
e e	1251	CGACCTCCCC	GAATCCCACC	TGAACGAGCT	GGCGGCCAGG	GGCCTGCCCA
55	1301	CGCCCGTGGT	TCTGGAATTC	GACAGCGAAT	TCGAGATGCT	GTTGGCCTTC
	1351	ATGACCCTTC	G TGAAACAGTA	CGGCCCCGAG	TTCGTGACCO	GGTACAACAT
60	1401	CATCAACTTC	GACTGGCCCT	TCTTGCTGGC	CAAGCTGAC	G GACATTTACA
	1451	AGGTCCCCCT	T GGACGGGTAC	GGCCGCATGA	ACGGCCGGG	G CGTGTTTCGC
6 5	1501	GTGTGGGAC	A TAGGCCAGAG	CCACTTCCAG	AAGCGCAGC	A AGATAAAGGT
65	1551	GAACGGCAT	G GTGAACATCG	ACATGTACGG	GATTATAAC	C GACAAGATCA

	1601	AGCTCTCGAG	CTACAAGCTC	AACGCCGTGG	CCGAAGCCGT	CCTGAAGGAC
5	1651	AAGAAGAAGG	ACCTGAGCTA	TCGCGACATC	CCCGCCTACT	ACGCCGCCGG
	1701	GCCCGCGCAA	CGCGGGGTGA	TCGGCGAGTA	CTGCATACAG	GATTCCCTGC
	1751	TGGTGGGCCA	GCTGTTTTT	AAGTTTTTGC	CCCATCTGGA	GCTCTCGGCC
10	1801	GTCGCGCGCT	TGGCGGGTAT	TAACATCACC	CGCACCATCT	ACGACGGCCA
	1851	GCAGATCCGC	GTCTTTACGT	GCCTGCTGCG	CCTGGCCGAC	CAGAAGGGCT
15	1901	TTATTCTGCC	GGACACCCAG	GGGCGATTTA	GGGGCGGCGG	GGGGGAGGCG
13	1951	CCCAAGCGTC	CGGCCGCAGC	CCGGGAGGAC	GAGGAGCGGC	CAGAGGAGGA
	2001	GGGGGAGGAC	GAGGACGAAC	GCGAGGAGGG	CGGGGGCGAG	CGGGAGCCGG
20	2051	AGGGCGCGCG	GEAGACCGCC	GGCCGGCACG	TGGGGTACCA	GGGGGCCAGG
	2101	GTCCTTGACC	CCACTTCCGG	GTTTCATGTG	AACCCCGTGG	TGGTGTTCGA
25	2151	CTTTGCCAGC	CTGTACCCCA	GCATCATCCA	GGCCCACAAC	CŢGTGCTTCA
23	2201	GCACGCTCTC	CCTGAGGGCC	GACGCAGTGG	CGCACCTGGA	GGCGGGÇAAG
	2251	GACTACCTGG	AGATCGAGGT	GGGGGGGCGA	CGGCTGTTCT	TCGTCAAGGC
30	2301	TCACGTGCGA	GAGAGCCTCC	TCAGCATCCT	CCTGCGGGAC	TGGCTCGCCA
	2351	TGCGAAAGCA	GATCCGCTCG	CGGATTCCCC	AGAGCAGCCC	CGAGGAGGCC
35	2401	GTGCTCCTGG	ACAAGCAGCA	GGCCGCCATC	AAGGTCGTGT	GTAACTCGGT
<i>33</i>	2451	TTACGGGTTC	ACGGGAGCGC	AGCACGGACT	CCTGCCGTGC	CTGCACGTTG
	2501	CCGCGACGGT	GACGACCATC	GGCCGCGAGA	TGCTGCTCGC	GACCCGCGAG
40	2551	TACGTCCACG	CGCGCTGGGC	GGCCTTCGAA	CAGCTCCTGG	CCGATTTCCC
	2601	GGAGGCGGCC	GACATGCGCG	CCCCGGGCC	CTATTCCATG	CGCATCATCT
45	2651	ACGGGGACAC	GGACTCCATC	TTTGTGCTGT	GCCGCGGCCT	CACGGCCGCC
73	2701	GGGCTGACGG	CCGTGGGCGA	CAAGATGGCG	AGCCACATCT	CGCGCGCGCT
	2751	GTTTCTGTCC	CCCATCAAAC	TCGAGTGCGA	AAAGACGTTC	ACCAAGCTGC
50	2801	TGCTGATCGC	CAAGAAAAAG	TACATCGGCG	TCATCTACGG	GGGTAAGATG
	2851	CTCATCAAGG	GCGTGGATCT	GGTGCGCAAA	AACAACTGCG	CGTTTATCAA
55	2901	CCGCACCTCC	AGGGCCCTGG	TCGACCTGCT	GTTTTACGAC	GATACCGTAT
33	2951	CCGGAGCGGC	CGCCGCGTT	A GCCGAGCGCC	CCGCAGAGGA	GTGGCTGGCG
	3001	CGACCCCTGC	CCGAGGGAC	GCAGGCGTTC	GGGGCCGTCC	TCGTAGACGC
60	3051	CCATCGGCGC	ATCACCGACC	CGGAGAGGGA	CATCCAGGAC	TTTGTCCTCA
	3101	CCGCCGAACT	' GAGCAGACA	C CCGCGCGCGT	ACACCAACAA	GCGCCTGGCC
65	3151	CACCTGACGG	TGTATTACA	A GCTCATGGCC	CGCCGCGCGC	AGGTCCCGTC
00	3201	CATCAAGGAC	CGGATCCCG	r acgtgatcgt	GGCCCAGACC	CGCGAGGTAG

	WO 02/06513					PCT/US01/16525	
	3251	AGGAGACGGT	CGCGCGGCTG	GCCGCCCTCC	GCGAGCTCGA	CGCCGCCGCC	
	3301	CCAGGGGACG	AGCCCGCCCC	CCCCGCGGCC	CTGCCCTCCC		
5	3351	CCCCCGGGAG	ACGCCGTTGC	ATGCCGACCC	CCCGGGAGGC	GCGTCCAAGC	
	3401	CCCGCAAGCT	GCTGGTGTCC	GAGCTGGCCG	AGGATCCCGC	ATACGCCATT	
10	3451	GCCCACGGCG	TCGCCCTGAA	CACGGACTAT	TACTTCTCCC	ACCTGTTGGG	
10	3501	GGCGGCGTGC	GTGACATTCA	AGGCCCTGTT	TGGGAATAAC	GCCAAGATCA	
	3551	CCGAGAGTCT	GTTAAAAAGG	TTTATTCCCG	AAGTGTGGCA	CCCCCGGAC	
15	3601	GACGTGGCCG	CGCGGCTCCG	GGCCGCAGGG	TTCGGGGCGG	TGGGTGCCGG	
	3651	CGCTACGGCG	GAGGAAACTC	GTCGAATGTT	GCATAGAGCC	TTTGATACTC	
	3701	TAGCATGA					

SEQ.ID.NO. 8 Amino acid sequence of DNA polymerase for HSV1-F-M1

5	1 MFSGGGGPLS PGGKSAARAA SGFFAPAGPR GAGRGPPPCL RQNFYNPYLA
	51 PVGTQQKPTG PTQRHTYYSE CDEFRFIAPR VLDEDAPPEK RAGVHDGHLK
	101 RAPKVYCGGD ERDVLRVGSG GFWPRRSRLW GGVDHAPAGF NPTVTVFHVY
10	151 DILENVEHAY GMRAAQFHAR FMDAITPTGT VITLLGLTPE GHRVAVHVYG
	201 TRQYFYMNKE EVDRHLQCRA PRDLCERMAA ALRESPGASF RGISADHFEA
	251 EVVERTDVYY YETRPALFYR VYVRSGRVLS YLCDNFCPAI KKYEGGVDAT
15	301 TRFILDNPGF VTFGWYRLKP GRNNTLAQPR APMAFGTSSD VEFNCTADNL
	351 AIEGGMSDLP AYKLMCFDIE CKAGGEDELA FPVAGHPEDL VIQISCLLYD
20	401 LSTTALEHVL LFSLGSCDLP ESHLNELAAR GLPTPVVLEF DSEFEMLLAF
20	451 MTLVKQYGPE FVTGYNIINF DWPFLLAKLT DIYKVPLDGY GRMNGRGVFR
	501 VWDIGQSHFQ KRSKIKVNGM VNIDMYGIIT DKIKLSSYKL NAVAEAVLKD
25	551 KKKDLSYRDI PAYYAAGPAQ RGVIGEYCIQ DSLLVGQLFF KFLPHLELSA
	601 VARLAGINIT RTTYDGQQIR VFTCLLRLAD QKGFILPDTQ GRFRGGGGEA
30	651 PKRPAAARED EERPEEEGED EDEREEGGGE REPEGARETA GRHVGYQGAR
30	701 VLDPTSGFHV NPVVVFDFAS LYPSIIQAHN LCFSTLSLRA DAVAHLEAGK
	751 DYLEIEVGGR RLFFVKAHVR ESLLSILLRD WLAMRKQIRS RIPQSSPEEA
35	801 VLLDKQQAAI KVVCNSVYGF TGAQHGLLPC LHVAATVTTI GREMLLATRE
	851 YVHARWAAFE QLLADFPEAA DMRAPGPYSM RIIYGDTDSI FVLCRGLTAA
40	901 GLTAVGDKMA SHISRALFLS PIKLECEKTF TKLLLIAKKK YIGVIYGGKM
40	951 LIKGVDLVRK NNCAFINRTS RALVDLLFYD DTVSGAAAAL AERPAEEWLA
	1001 RPLPEGLQAF GAVLVDAHRR ITDPERDIQD FVLTAELSRH PRAYTNKRLA
45	1051 HLTVYYKLMA RRAQVPSIKD RIPYVIVAQT REVEETVARL AALRELDAAA
	1101 PGDEPAPPAA LPSPAKRPRE TPLHADPPGG ASKPRKLLVS ELAEDPAYAI
50	1151 AHGVALNTDY YFSHLLGAAC VTFKALFGNN AKITESLLKR FIPEVWHPPD
30	1201 DVAARLRAAG FGAVGAGATA EETRRMLHRA FDTLA*

SEQ.ID.NO. 9 DNA sequence of HSV polymerase gene for HSV1-DJL-M1

1 ATGTTTTCCG GTGGCGGCGG CCCGCTGTCC CCCGGAGGAA AGTCGGCGGC 51 CAGGGCGCG TCCGGGTTTT TTGCGCCCGC CGGCCCTCGC GGAGCCGGCC 5 101 GGGGACCCCC GCCTTGTTTG AGGCAAAACT TTTACAACCC CTACCTCGCC 151 CCAGTCGGGA CGCAACAGAA GCCGACCGGG CCAACCCAGC GCCATACGTA 10 201 CTATAGCGAA TGCGATGAAT TTCGATTCAT CGCCCCGCGG GTGCTGGACG 251 AGGATGCCCC CCCGGAGAAG CGCGCCGGGG TGCACGACGG TCACCTCAAG 301 CGCGCCCCCA AGGTGTACTG CGGGGGGGAC GAGCGCGACG TCCTCCGCGT 15 351 CGGGTCGGGC GGCTTCTGGC CGCGGCGCTC GCGCCTGTGG GGCGGCGTGG 401 ACCACGCCCC GGCGGGGTTC AACCCCACCG TCACCGTCTT TCACGTGTAT 20 451 GACATCCTGG AGAACGTGGA GCACGCGTAC GGCATGCGCG CGGCCCAGTT 501 CCACGCGCGG TTTATGGACG CCATCACACC GACGGGGACC GTCATCACGC 551 TCCTGGGCCT GACTCCGGAA GGCCACCGGG TGGCCGTTCA CGTTTACGGC 25 601 ACGCGGCAGT ACTTTTACAT GAACAAGGAG GAGGTTGACA GGCACCTACA 651 ATGCCGCGCC CCACGAGATC TCTGCGAGCG CATGGCCGCG GCCCTGCGCG 30 701 AGTCCCCGGG CGCGTCGTTC CGCGGCATCT CCGCGGACCA CTTCGAGGCG 751 GAGGTGGTGG AGCGCACCGA CGTGTACTAC TACGAGACGC GCCCCGCTCT 801 GTTTTACCGC GTCTACGTCC GAAGCGGGCG CGTGCTGTCG TACCTGTGCG 35 851 ACAACTTCTG CCCGGCCATC AAGAAGTACG AGGGTGGGGT CGACGCCACC 901 ACCCGGTTCA TCCTGGACAA CCCCGGGTTC GTCACCTTCG GCTGGTACCG 40 951 TCTCAAACCG GGCCGGAACA ACACGCTAGC CCAGCCGCGG GCCCCGATGG 1001 CCTTCGGGAC ATCCAGCGAT GTCGAGTTTA ACTGTACGGC GGACAACCTG 1051 GCCATCGAGG GGGGCATGAG CGACCTACCG GCATACAAGC TCATGTGCTT 45 1101 CGATATCGAA TGCAAGGCGG GGGGGGAGGA CGAGCTGGCC TTTCCGGTGG 1151 CCGGGCACCC GGAGGACCTG GTCATCCAGA TATCCTGTCT GCTCTACGAC 50 1201 CTGTCCACCA CCGCCCTGGA GCACGTCCTC CTGTTTTCGC TCGGTTCCTG 1251 CGACCTCCCC GAATCCCACC TGAACGAGCT GGCGGCCAGG GGCCTGCCCA 1301 CGCCCGTGGT TCTGGAATTC GACAGCGAAT TCGAGATGCT GTTGGCCTTC 55 1351 ATGACCCTTG TGAAACAGTA CGGCCCCGAG TTCGTGACCG GGTACAACAT 1401 AATCAACTTC GACTGGCCCT TCTTGCTGGC CAAGCTGACG GACATTTACA

5

10

15

20

25

30

35

40

45

50

55

60

1451 AGGTCCCCCT GGACGGGTAC GGCCGCATGA ACGGCCGGGG CGTGTTTCGC 1501 GTGTGGGACA TAGGCCAGAG CCACTTCCAG AAGCGCAGCA AGATAAAGGT 1551 GAACGCCATG GTGAACATCG ACATGTACGG GATTATAACC GACAAGATCA 1601 AGCTCTCGAG CTACAAGCTC AACGCCGTGG CCGAAGCCGT CCTGAAGGAC 1651 AAGAAGAAGG ACCTGAGCTA TCGCGACATC CCCACCTACT ACGCCGCCGG 1701 GCCCGCGCAA CGCGGGGTGA TCGGCGAGTA CTGCATACAG GATTCCCTGC 1751 TGGTGGGCCA GCTGTTTTTT AAGTTTTTTGC CCCATCTGGA GCTCTCGGCC 1801 GTCGCGCGCT TGGCGGGTAT TAACATCACC CGCACCATCT ACGACGCCA 1851 GCAGATCCGC GTCTTTACGT GCCTGCTGCG CCTGGCCGAC CAGAAGGGCT 1901 TTATTCTGCC GGACACCCAG GGGCGATTTA GGGGCGCCGG GGGGGAGGCG 1951 CCCAAGCGTC CGGCCGCAGC CCGGGAGGAC GAGGAGCGC CAGAGGAGGA 2001 GGGGGAGGAC GAGAACGAAC GCGAGGAGGG CGGGGGCGAG CGGGAGCCGG 2051 AGGGCGCGCGGGAGACCGCC GGCCGGCACG TGGGGTACCA GGGGGCCAGG 2101 GTCCTTGACC CCACTTCCGG GTTTCACGTG AACCCCGTGG TGGTGTTCGA 2151 CTTTGCCAGC CTGTACCCCA GCATCATCCA GGCCCACAAC CTGTGCTTCA 2201 GCACGCTCTC CCTGAGGGCC GACGCAGTGG CGCACCTGGA GGCGGGCAAG 2251 GACTACCTGG AGATCGAGGT GGGGGGGGGGA CGGCTGTTCT TCGTCAAGGC 2301 TCACGTGCGA GAGAGCCTCC TCAGCATCCT CCTGCGGGAC TGGCTCGCCA 2351 TGCGAAAGCA GATCCGCTCG CGGATTCCCC AGAGCAGCCC CGAGGAGGCC 2401 GTGCTCCTGG ACAAGCAGCA GGCCGCCATC AAGGTCGTGT GTAACTCGGT 2451 TTACGGGTTC ACGGGAGCGC AGCACGGACT CCTGCCGTGC CTGCACGTTG 2501 CCGCGACGGT GACGACCATC GGCCGCGAGA TGCTGCTCGC GACCCGCGAG 2551 TACGTCCACG CGCGCTGGGC GGCCTTCGAA CAGCTCCTGG CCGATTTCCC 2601 GGAGGCGCC GACATGCGCG CCCCCGGGCC CTATTCCATG CGCATCATCT 2651 ACGGGGACAC GGACTCCATA TTTGTGCTGT GCCGCGGCCT CACGGCCGCC 2701 GGGCTGACGG CCGTGGGCGA CAAGATGGCG AGCCACATCT CGCGCGCGCT 2751 GTTTCTGCCC CCCATCAAAC TCGAGTGCGA AAAGACGTTC ACCAAGCTGC 2801 TGCTGATCGC CAAGAAAAAG TACATCGGCG TCATCTACGG GGGTAAGATG 2851 CTCATCAAGG GCGTGGATCT GGTGCGCAAA AACAACTGCG CGTTTATCAA 2901 CCGCACCTCC AGGGCCCTGG TCGACCTGCT GTTTTACGAC GATACCGTAT

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25

30

2951 CCGGAGCGC CGCCGCGTTA GCCGAGCGCC CCGCAGAGGA GTGGCTGGCG 3001 CGACCCCTGC CCGAGGGACT GCAGGCGTTC GGGGCCGTCC TCGTAGACGC 3051 CCATCGGCGC ATCACCGACC CGGAGAGGGA CATCCAGGAC TTTGTTCTCA 3101 CCGCCGAACT GAGCAGACAC CCGCGCGCGT ACACCAACAA GCGCCTGGCC 3151 CACCTGACGG TGTATTACAA GCTCATGGCC CGCCGCGCGC AGGTCCCGTC 3201 CATCAAGGAC CGGATCCCGT ACGTGATCGT GGCCCAGACC CGCGAGGTAG 3251 AGGAGACGGT CGCGCGGCTG GCCGCCCTCC GCGAGCTAGA CGCCGCCGCC 3301 CCAGGGGACG AGCCCGCCCC CCCCGCGGCC CTGCCCTCCC CGGCCAAGCG 3351 CCCCCGGGAG ACGCCGTCGC CTGCCGACCC CCCGGGAGGC GCGTCCAAGC 3401 CCCGCAAGCT GCTGGTGTCC GAGCTGGCCG AGGATCCCGC ATACGCCATT 3451 GCCCACGGCG TCGCCCTGAA CACGGACTAT TACTTCTCCC ACCTGTTGGG 3501 GGCGGCGTGC GTGACATTCA AGGCCCTGTT TGGGAATAAC GCCAAGATCA 3551 CCGAGAGTCT GTTAAAAAGG TTTATTCCCG AAGTGTGGCA CCCCCCGGAC 3601 GACGTGGCCG CGCGGCTCCG GACCGCAGGG TTCGGGGCGG TGGGTGCCGG 3651 CGCTACGGCG GAGGAAACTC GTCGAATGTT GCATAGAGCC TTTGATACTC 3701 TAGCATGA

SEQ.ID.NO. 10 Amino acid sequence of DNA polymerase for HSV1-DJL-M1

	1 MFSGGGGPLS PGGKSAARAA SGFFAPAGPR GAGRGPPPCL RQNFYNPYLA
5	51 PVGTQQKPTG PTQRHTYYSE CDEFRFIAPR VLDEDAPPEK RAGVHDGHLK
	101 RAPKVYCGGD ERDVLRVGSG GFWPRRSRLW GGVDHAPAGF NPTVTVFHVY
10	151 DILENVEHAY GMRAAQFHAR FMDAITPTGT VITLLGLTPE GHRVAVHVYG
10	201 TRQYFYMNKE EVDRHLQCRA PRDLCERMAA ALRESPGASF RGISADHFEA
	251 EVVERTDVYY YETRPALFYR VYVRSGRVLS YLCDNFCPAI KKYEGGVDAT
15	301 TRFILDNPGF VTFGWYRLKP GRNNTLAQPR APMAFGTSSD VEFNCTADNL
	351 AIEGGMSDLP AYKLMCFDIE CKAGGEDELA FPVAGHPEDL VIQISCLLYD
20	401 LSTTALEHVL LFSLGSCDLP ESHLNELAAR GLPTPVVLEF DSEFEMLLAF
20	451 MTLVKQYGPE FVTGYNIINF DWPFLLAKLT DIYKVPLDGY GRMNGRGVFR
	501 VWDIGQSHFQ KRSKIKVNGM VNIDMYGIIT DKIKLSSYKL NAVAEAVLKD
25	551 KKKDLSYRDI PTYYAAGPAQ RGVIGEYCIQ DSLLVGQLFF KFLPHLELSA
	601 VARLAGINIT RTIYDGQQIR VFTCLLRLAD QKGFILPDTQ GRFRGAGGEA
30	651 PKRPAAARED EERPEEEGED ENEREEGGGE REPEGARETA GRHVGYQGAR
50	701 VLDPTSGFHV NPVVVFDFAS LYPSIIQAHN LCFSTLSLRA DAVAHLEAGK
	751 DYLEIEVGGR RLFFVKAHVR ESLLSILLRD WLAMRKQIRS RIPQSSPEEA
35	801 VLLDKQQAAI KVVCNSVYGF TGAQHGLLPC LHVAATVTTI GREMLLATRE
	851 YVHARWAAFE QLLADFPEAA DMRAPGPYSM RIIYGDTDSI FVLCRGLTAA
40	901 GLTAVGDKMA SHISRALFLP PIKLECEKTF TKLLLIAKKK YIGVIYGGKM
40	951 LIKGVDLVRK NNCAFINRTS RALVDLLFYD DTVSGAAAAL AERPAEEWLA
	1001 RPLPEGLQAF GAVLVDAHRR ITDPERDIQD FVLTAELSRH PRAYTNKRLA
45	1051 HLTVYYKLMA RRAQVPSIKD RIPYVIVAQT REVEETVARL AALRELDAAA
	1101 PGDEPAPPAA LPSPAKRPRE TPSPADPPGG ASKPRKLLVS ELAEDPAYAI
50	1151 AHGVALNTDY YFSHLLGAAC VTFKALFGNN AKITESLLKR FIPEVWHPPD
50	1201 DVAARLRTAG FGAVGAGATA EETRRMLHRA FDTLA*

SEQ.ID.NO. 11 DNA sequence of DNA polymerase gene for HMCV-AD169-M1

1 ATGTTTTTCA ACCCGTATCT GAGCGGCGGC GTGACCGGCG GTGCGGTCGC 51 GGGTGGCCGG CGTCAGCGTT CGCAGCCCGG CTCCGCGCAG GGCTCGGGCA 5 101 AGCGGCCGCC ACAGAAACAG TTTTTGCAGA TCGTGCCGCG AGGTGTCATG 151 TTCGACGGTC AGACGGGGTT GATCAAGCAT AAGACGGGAC GGCTGCCTCT 10 201 CATGTTCTAT CGAGAGATTA AACATTTGTT GAGTCATGAC ATGGTTTGGC 251 CGTGTCCTTG GCGCGAGACC CTGGTGGGTC GCGTGGTGGG ACCTATTCGT 301 TTTCACACCT ACGATCAGAC GGACGCCGTG CTCTTCTTCG ACTCGCCCGA 15 351 AAACGTGTCG CCGCGCTATC GTCAGCATCT GGTGCCTTCG GGGAACGTGT 401 TGCGTTTCTT CGGGGCCACA GAACACGGCT ACAGTATCTG CGTCAACGTT 20 451 TTCGGGCAGC GCAGCTACTT TTACTGTGAG TACAGCGACA CCGATAGGCT 501 GCGTGAGGTC ATTGCCAGCG TGGGCGAACT AGTGCCCGAA CCGCGGACGC 551 CATACGCCGT GTCTGTCACG CCGGCCACCA AGACCTCCAT CTATGGGTAC 25 601 GGGACGCGAC CCGTGCCCGA TTTGCAGTGT GTGTCTATCA GCAACTGGAC 651 CATGGCCAGA AAAATCGGCG AGTATCTGCT GGAGCAGGGT TTTCCCGTGT 30 701 ACGAGGTCCG TGTGGATCCG CTGACGCGTT TGGTCATCGA TCGGCGGATC 751 ACCACGTTCG GCTGGTGCTC CGTGAATCGT TACGACTGGC GGCAGCAGGG 801 TCGCGCGTCG ACTTGTGATA TCGAGGTAGA CTGCGATGTC TCTGACCTGG 35 851 TGGCTGTGCC CGACGACAGC TCGTGGCCGC GCTATCGATG CCTGTCCTTC 901 GATATCGAGT GCATGAGCGG CGAGGGTGGT TTTCCCTGCG CCGAGAAGTC 40 951 CGATGACATT GTCATTCAGA TCTCGTGCGT GTGCTACGAG ACGGGGGAA 1001 ACACCGCCGT GGATCAGGGG ATCCCAAACG GGAACGATGG TCGGGGCTGC 1051 ACTTCGGAGG GTGTGATCTT TGGGCACTCG GGTCTTCATC TCTTTACGAT 45 1101 CGGCACCTGC GGGCAGGTGG GCCCAGACGT GGACGTCTAC GAGTTCCCTT 1151 CCGAATACGA GCTGCTGCTG GGCTTTATGC TTTTCTTTCA ACGGTACGCG 50 1201 CCGGCCTTTG TGACCGGTTA CAACATCAAC TCTTTTGACT TGAAGTACAT 1251 CCTCACGCGT CTCGAGTACC TGTATAAGGT GGACTCGCAG CGCTTCTGCA 1301 AGTTGCCTAC GGCGCAGGGC GGCCGTTTCT TTTTACACAG CCCCGCCGTG 55 1351 GGTTTTAAGC GGCAGTACGC CGCCGCTTTT CCCTCGGCTT CTCACAACAA 1401 TCCGGCCAGC ACGCCGCCA CCAAGGTGTA TATTGCGGGT TCGGTGGTTA

1451 TCGACATGTA CCCTGTATGC ATGGCCAAGA CTAACTCGCC CAACTATAAG 1501 CTCAACACTA TGGCCGAGCT TTACCTGCGG CAACGCAAGG ATGACCTGTC • 5 1551 TTACAAGGAC ATCCCGCGTT GTTTCGTGGC TAATGCCGAG GGCCGCCCC 1601 AGGTAGGCCG TTACTGTCTG CAGGACGCCG TATTGGTGCG CGATCTGTTC 1651 AACACCATTA ATTTTCACTA CGAGGCCGGG GCCATCGCGC GGCTGGCTAA 10 1701 AATTCCGTTG CGGCGTGTCA TCTTTGACGG ACAGCAGATC CGTATCTACA 1751 CCTCGCTGCT GGACGAGTGC GCCTGCCGCG ATTTTATCCT GCCCAACCAC 15 1801 TACAGCAAAG GTACGACGGT GCCCGAAACG AATAGCGTTG CTGTGTCACC 1851 TAACGCTGCT ATCATCTCTA CCGCCGCTGT GCCCGGCGAC GCGGGTTCTG 1901 TGGCGGCTAT GTTTCAGATG TCGCCGCCCT TGCAATCTGC GCCGTCCAGT 20 1951 CAGGACGCG TTTCACCCGG CTCCGGCAGT AACAGTAGTA GCAGCGTCGG 2001 CGTTTTCAGC GTCGGCTCCG GCAGTAGTGG CGGCGTCGGC GTTTCCAACG 25 2051 ACAATCACGG CGCCGGCGGT ACTGCGGCGG TTTCGTACCA GGGCGCCACG 2101 GTGTTTGAGC CCGAGGTGGG TTACTACAAC GACCCCGTGG CCGTGTTCGA 2151 CTTTGCCAGC CTCTACCCTT CCATCATCAT GGCCCACAAC CTCTGCTACT 30 2201 CCACCCTGCT GGTGCCGGGT GGCGAGTACC CTGTGGACCC CGCCGACGTA 2251 TACAGCGTCA CGCTAGAGAA CGGCGTGACC CACCGCTTTG TGCGTGCTTC 35 2301 GGTGCGCGTC TCGGTGCTCT CGGAACTGCT CAACAAGTGG GTTTCGCAGC 2351 GGCGTGCCGT GCGCGAATGC ATGCGCGAGT GTCAAGACCC TGTGCGCCGT 2401 ATGCTGCTCG ACAAGGAACA GATGGCGCTC AAAGTAACGT GCAACGCTTT 40 2451 CTACGGTTTT ACCGGCGCGC TGAACGGTAT GATGCCGTGT CTGCCCATCG 2501 CCGCCAGCAT CACGCGCATC GGTCGCGACA TGCTAGAGCG CACGGCGCGG 45 2551 TTCATCAAAG ACAACTTTTC AGAGCCGTGT TTTTTGCACA ATTTTTTAA 2601 TCAGGAAGAC TATGTAGTGG GAACGCGGGA GGGGGATTCG GAGGAGAGCA 2651 GCGCGTTACC GGAGGGGCTC GAAACATCGT CAGGGGGCTC GAACGAACGG 50 2701 CGGGTGGAGG CGCGGGTCAT CTACGGGGAC ACGGACAGCG TGTTTGTCCG 2751 CTTTCGTGGC CTGACGCCGC AGGCTCTGGT GGCGCGTGGG CCCAGCCTGG 55 2801 CGCACTACGT GACGGCCTGT CTTTTTGTGG AGCCCGTCAA GCTGGAGTTT 2851 GAAAAGGTCT TCGTCTCTCT TATGATGATC TGCAAGAAAC GTTACATCGG 2901 CAAAGTGGAG GGCGCCTCGG GTCTGAGCAT GAAGGGCGTG GATCTGGTGC 60

2951 GCAAGACGGC CTGCGAGTTC GTCAAGGGCG TCACGCGTGA CGTCCTCTCG 3001 CTGCTCTTTG AGGATCGCGA GGTCTCGGAA GCAGCCGTGC GCCTGTCGCG 5 3051 CCTCTCACTC GATGAAGTCA AGAAGTACGG CGTGCCACGC GGTTTCTGGC 3101 GTATCTTACG CCGCTTGGTG CAGGCCCGCG ACGATCTGTA CCTGCACCGT 3151 GTGCGTGTCG AGGACCTGGT GCTTTCGTCG GTGCTCTCTA AGGACATCTC 10 3201 GCTGTACCGT CAATCTAACC TGCCGCACAT TGCCGTCATT AAGCGATTGG 3251 CGGCCCGTTC TGAGGAGCTA CCCTCGGTCG GGGATCGGGT CTTTTACGTT 15 3301 CTGACGGCGC CCGGTGTCCG GACGGCGCCG CAGGGTTCCT CCGACAACGG 3351 TGATTCTGTA ACCGCCGGCG TGGTTTCCCG GTCGGACGCG ATTGATGGCA 3401 CGGACGACGA CGCTGACGGC GGCGGGGTAG AGGAGAGCAA CAGGAGAGGA 20 3451 GGAGAGCCGG CAAAGAAGAG GGCGCGGAAA CCACCGTCGG CCGTGTGCAA 3501 CTACGAGGTA GCCGAAGATC.CGAGCTACGT GCGCGAGCAC GGCGTGCCCA 25 3551 TTCACGCCGA CAAGTACTTT GAGCAGGTTC TCAAGGCTGT AACTAACGTG 3601 CTGTCGCCCG TCTTTCCCGG CGCGAAACC GCGCGCAAGG ACAAGTTTTT 3651 GCACATGGTG CTGCCGCGGC GCTTGCACTT GGAGCCGGCT TTTCTGCCGT 30 3701 ACAGTGTCAA GGCGCACGAA TGCTGTTGA

SEQ.ID.NO.12 Amino acid sequence of DNA polymerase for HCMV-AD169-M1

5	1 MFFNPYLSGG VTGGAVAGGR RQRSQPGSAQ GSGKRPPQKQ FLQIVPRGVM
	51 FDGQTGLIKH KTGRLPLMFY REIKHLLSHD MVWPCPWRET LVGRVVGPIR
	101 FHTYDQTDAV LFFDSPENVS PRYRQHLVPS GNVLRFFGAT EHGYSICVNV
10	151 FGQRSYFYCE YSDTDRLREV IASVGELVPE PRTPYAVSVT PATKTSIYGY
	201 GTRPVPDLQC VSISNWTMAR KIGEYLLEQG FPVYEVRVDP LTRLVIDRRI
15	251 TTFGWCSVNR YDWRQQGRAS TCDIEVDCDV SDLVAVPDDS SWPRYRCLSF
15	301 DIECMSGEGG FPCAEKSDDI VIQISCVCYE TGGNTAVDQG IPNGNDGRGC
	351 TSEGVIFGHS GLHLFTIGTC GQVGPDVDVY EFPSEYELLL GFMLFFQRYA
20	401 PAFVTGYNIN SFDLKYILTR LEYLYKVDSQ RFCKLPTAQG GRFFLHSPAV
	451 GFKRQYAAAF PSASHNNPAS TAATKVYIAG SVVIDMYPVC MAKTNSPNYK
25	501 LNTMAELYLR QRKDDLSYKD IPRCFVANAE GRAQVGRYCL QDAVLVRDLF
<i></i>	551 NTINFHYEAG AIARLAKIPL RRVIFDGQQI RIYTSLLDEC ACRDFILPNH
•	601 YSKGTTVPET NSVAVSPNAA IISTAAVPGD AGSVAAMFQM SPPLQSAPSS
30	651 QDGVSPGSGS NSSSSVGVFS VGSGSSGGVG VSNDNHGAGG TAAVSYQGAT
	701 VFEPEVGYYN DPVAVFDFAS LYPSIIMAHN LCYSTLLVPG GEYPVDPADV
35	751 YSVTLENGVT HRFVRASVRV SVLSELLNKW VSQRRAVREC MRECQDPVRR
	801 MLLDKEQMAL KVTCNAFYGF TGALNGMMPC LPIAASITRI GRDMLERTAR
	851 FIKDNFSEPC FLHNFFNQED YVVGTREGDS EESSALPEGL ETSSGGSNER
40	901 RVEARVIYGD TDSVFVRFRG LTPQALVARG PSLAHYVTAC LFVEPVKLEF
	951 EKVFVSLMMI CKKRYIGKVE GASGLSMKGV DLVRKTACEF VKGVTRDVLS
45	1001 LLFEDREVSE AAVRLSRLSL DEVKKYGVPR GFWRILRRLV QARDDLYLHR
	1051 VRVEDLVLSS VLSKDISLYR QSNLPHIAVI KRLAARSEEL PSVGDRVFYV
	1101 LTAPGVRTAP QGSSDNGDSV TAGVVSRSDA IDGTDDDADG GGVEESNRRG
50	1151 GEPAKKRARK PPSAVCNYEV AEDPSYVREH GVPIHADKYF EQVLKAVTNV
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Figure 6
SEQ.ID.NO.13 Amino acid sequence of DNA polymerase for HCMV-AD169

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20	401 PAFVTGYNIN SFDLKYILTR LEYLYKVDSQ RFCKLPTAQG GRFFLHSPAV
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25	501 LNTMAELYLR QRKDDLSYKD IPRCFVANAE GRAQVGRYCL QDAVLVRDLF
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SEQUENCE LISTING

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Ile 305	Leu	Asp	Asn	Pro	Gly 310	Phe	Val	Thr	Phe	Gly 315	Trp	Tyr	Arg	Leu	Lys 320
Pro	Gly	Arg	Gly	Asn 325	Ala	Pro	Ala	Gln	Pro 330	Arg	Pro	Pro	Thr	Ala 335	Phe
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				405					410					Leu 415	
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Leu	Pro	Ala 435	Pro	Val	Val	Leu	Glu 440	Phe	Asp	Ser	Glu	Phe 445	Glu	Met	Leu
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Gly 465	Tyr	Asn	Ile	Ile	Asn 470	Phe	Asp	Trp	Pro	Phe 475	Val	Leu	Thr	Lys	Leu 480

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Val Val Cys Asn Ser Val Tyr Gly Phe Thr Gly Ala Gln His Gly Leu 820 825 830

- Leu Pro Cys Leu His Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu 835 840 845
- Met Leu Leu Ala Thr Arg Ala Tyr Val His Ala Arg Trp Ala Glu Phe 850 855 860
- Asp Gln Leu Leu Ala Asp Phe Pro Glu Ala Ala Gly Met Arg Ala Pro 865 870 875 880
- Gly Pro Tyr Ser Met Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe 885 890 895
- Val Leu Cys Arg Gly Leu Thr Ala Ala Gly Leu Val Ala Met Gly Asp 900 905 910
- Lys Met Ala Ser His Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys 915 920 925
- Leu Glu Cys Glu Lys Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys 930 935 940
- Lys Tyr Ile Gly Val Ile Cys Gly Gly Lys Met Leu Ile Lys Gly Val 945 950 955 960
- Asp Leu Val Arg Lys Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg 965 970 975
- Ala Leu Val Asp Leu Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala 980 985 990
- Ala Ala Leu Ala Glu Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu 995 1000 1005
- Pro Glu Gly Leu Gln Ala Phe Gly Ala Val Leu Val Asp Ala His 1010 1015 1020
- Arg Arg Ile Thr Asp Pro Glu Arg Asp Ile Gln Asp Phe Val Leu 1025 1030 1035
- Thr Ala Glu Leu Ser Arg His Pro Arg Ala Tyr Thr Asn Lys Arg 1040 1045 1050
- Leu Ala His Leu Thr Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala 1055 1060 1065
- Gln Val Pro Ser Ile Lys Asp Arg Ile Pro Tyr Val Ile Val Ala 1070 1075 1080
- Gln Thr Arg Glu Val Glu Glu Thr Val Ala Arg Leu Ala Ala Leu 1085 1090 1095
- Arg Glu Leu Asp Ala Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro 1100 1105 1110 .
- Ala Ala Leu Pro Ser Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser 1115 1120 1125
- His Ala Asp Pro Pro Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu

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	Pro 1160	Leu	Asn	Thr	Asp	Tyr 1165	Tyr	Phe	Ser	His	Leu 1170	Leu	Gly	Ala	
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	Glu 1190	Ser	Leu	Leu	Lys	Arg 1195	Phe	Ile	Pro	Glu	Thr 1200	Trp	His	Pro	
	Asp 1205	Asp	Val	Ala	Ala	Arg 1210	Leu	Arg	Ala	Ala	Gly 1215	Phe	Gly	Pro	
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Leu Ala Pro Val Gly Thr Gln Gln Lys Pro Thr Gly Pro Thr Gln Arg 50 55 60

His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 75 80

Val Leu Asp Glu Asp Ala Pro Pro Glu Lys Arg Ala Gly Val His Asp 85 90 95

Gly His Leu Lys Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu Arg 100 105 110

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Arg	Val	Ala 195	Val	His	Val	Tyr	Gly 200	Thr	Arg	Gln	Tyr	Phe 205	Tyr	Met	Asn
Lys	Glu 210	Glu	Val	Asp	Arg	His 215	Leu	Gln	Cys	Arg	Ala 220	Pro	Arg	Asp	Leu
Суs 225	Glu	Arg	Met	Ala	Ala 230	Ala	Leu	Àrg	Glu	Ser 235	Pro	Gly	Āla	Ser	Phe 240
Arg	Gly	Ile	Ser	Ala 245	Asp	His	Phe	Glu	Ala 250	Glu	Val	Val	Glu	Arg 255	Thr
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Val	Arg	Ser 2 7 5	Gly	Arg	Val	Leu	Ser 280	Tyr	Leu	Cys	Asp	Asn 285	Phe	Cys	Pro
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	Gly	355					360					365			
	Glu 370					375					380				
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	Thr	435					440					445			
Ala	Phe 450	Met	Thr	Leu	Val	Lys 455	Gln	Tyr	Gly	Pro	Glu 460	Phe	Val	Thr	Gly

Tyr Asn Ile Ile Asn Phe Asp Trp Pro Phe Leu Leu Ala Lys Leu Thr Asp Ile Tyr Lys Val Pro Leu Asp Gly Tyr Gly Arg Met Asn Gly Arg Gly Val Phe Arg Val Trp Asp Ile Gly Gln Ser His Phe Gln Lys Arg Ser Lys Ile Lys Val Asn Gly Met Val Asn Ile Asp Met Tyr Gly Ile Ile Thr Asp Lys Ile Lys Leu Ser Ser Tyr Lys Leu Asn Ala Val Ala Glu Ala Val Leu Lys Asp Lys Lys Lys Asp Leu Ser Tyr Arg Asp Ile Pro Ala Tyr Tyr Ala Ala Gly Pro Ala Gln Arg Gly Val Ile Gly Glu Tyr Cys Ile Gln Asp Ser Leu Leu Val Gly Gln Leu Phe Phe Lys Phe Leu Pro His Leu Glu Leu Ser Ala Val Ala Arg Leu Ala Gly Ile Asn Ile Thr Arg Thr Ile Tyr Asp Gly Gln Gln Ile Arg Val Phe Thr Cys Leu Leu Arg Leu Ala Asp Gln Lys Gly Phe Ile Leu Pro Asp Thr Gln Gly Arg Phe Arg Gly Ala Gly Gly Glu Ala Pro Lys Arg Pro Ala Ala Ala Arg Glu Asp Glu Glu Arg Pro Glu Glu Glu Glu Glu Asp Glu Asp Glu Arg Glu Glu Gly Gly Glu Arg Glu Pro Glu Gly Ala Arg Glu Thr Ala Gly Arg His Val Gly Tyr Gln Gly Ala Arg Val Leu Asp Pro Thr Ser Gly Phe His Val Asn Pro Val Val Val Phe Asp Phe Ala Ser Leu Tyr Pro Ser Ile Ile Gln Ala His Asn Leu Cys Phe Ser Thr Leu Ser Leu Arg Ala Asp Ala Val Ala His Leu Glu Ala Gly Lys Asp Tyr Leu Glu Ile Glu Val Gly Gly Arg Arg Leu Phe Phe Val Lys Ala His Val Arg Glu Ser Leu Leu Ser Ile Leu Leu Arg Asp Trp Leu Ala Met Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala

. 17

- Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val Cys Asn Ser 805 810 815
- Val Tyr Gly Phe Thr Gly Ala Gln His Gly Leu Leu Pro Cys Leu His 820 825 830
- Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu Leu Ala Thr 835 840 845
- Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala 850 855 860
- Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met 865 870 875 880
- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Met Gly Asp Lys Met Ala Ser His
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- Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
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- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr 1040 1045 1050
- Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala 1085 1090 1095
- Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala Leu Pro Ser 1100 1105 1110
- Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser His Ala Asp Pro Pro

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Lys	Ala 1175	Leu	Phe	Gly	Asn	Asn 1180	Ala	Lys	Ile	Thr	Glu 1185	Ser	Leu	Leu	
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Leu	Ala 1235														
			s sin	mplex	¢										
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Pro Thr Pro Val Val Leu Glu Phe Asp Ser Glu Phe Glu Met Leu Leu

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Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala 785 790 795 800

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- Val Tyr Gly Phe Thr Gly Ala Gln His Gly Leu Leu Pro Cys Leu His 820 825 830
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- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 885 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Val Gly Asp Lys Met Ala Ser His
 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Ser Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys 945 950 955 960
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
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- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
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His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 75 80

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- Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala 850 855 860
- Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met 865 870 . 875 880
- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 885 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Val Gly Asp Lys Met Ala Ser His 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys 945 950 955 960
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990 ,
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 · 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr 1040 1045 1050
- Val Tyr Tyr Lys. Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr 'Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala

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Gľu	Asp 1145	Pro	Ala	Tyr	Ala	Ile 1150	Ala	His	Gly	Val	Ala 1155	Leu	Asn	Thr	
Asp	Tyr 1160	-	Phe	Ser	His	Leu 1165	Leu	Gly	Ala	Ala	Cys 1170	Val	Thr	Phe	
Ъγъ	Ala 1175		Phe	Ğİy	Asn	Asn 1180	Ala	Lys	Ile	Thr	Glu 1185	Ser	Leu	Leu	
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Ala	Arg 1205		Arg	Thr	Ala	Gly 1210		Gly	Ala	Val	Gly 1215	Ala	Gly	Ala	
Thr	Ala 1220		Glu	Thr	Arg	Arg 1225		Leu	His	Arg	Ala 1230	Phe	Asp	Thr	
Leu	Ala 1235														
	_	729 NA	s si	mple:	×	•									
-	0> 1		cccg	tatc	t ga	gcggc	ggc (gtga	ccgg	cg g	tgcgg	tcgc	ggg	tggccgg	r 60
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Ala Gly Gly Arg Arg Gln Arg Ser Gln Pro Gly Ser Ala Gln Gly Ser 20 25 30

Gly Lys Arg Pro Pro Gln Lys Gln Phe Leu Gln Ile Val Pro Arg Gly 35 40 45

Val Met Phe Asp Gly Gln Thr Gly Leu Ile Lys His Lys Thr Gly Arg 50 55 60

Leu Pro Leu Met Phe Tyr Arg Glu Ile Lys His Leu Leu Ser His Asp
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<212> PRT

<213> herpes simplex

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Ile Leu Thr Arg Leu Glu Tyr Leu Tyr Lys Val Asp Ser Gln Arg Phe Cys Lys Leu Pro Thr Ala Gln Gly Gly Arg Phe Phe Leu His Ser Pro Ala Val Gly Phe Lys Arg Gln Tyr Ala Ala Ala Phe Pro Ser Ala Ser His Asn Asn Pro Ala Ser Thr Ala Ala Thr Lys Val Tyr Ile Ala Gly Ser Val Val Ile Asp Met Tyr Pro Val Cys Met Ala Lys Thr Asn Ser Pro Asn Tyr Lys Leu Asn Thr Met Ala Glu Leu Tyr Leu Arg Gln Arg Lys Asp Asp Leu Ser Tyr Lys Asp Ile Pro Arg Cys Phe Val Ala Asn Ala Glu Gly Arg Ala Gln Val Gly Arg Tyr Cys Leu Gln Asp Ala Val Leu Val Arg Asp Leu Phe Asn Thr Ile Asn Phe His Tyr Glu Ala Gly Ala Ile Ala Arg Leu Ala Lys Ile Pro Leu Arg Arg Val Ile Phe Asp Gly Gln Gln Ile Arg Ile Tyr Thr Ser Leu Leu Asp Glu Cys Ala Cys Arg Asp Phe Ile Leu Pro Asn His Tyr Ser Lys Gly Thr Thr Val Pro Glu Thr Asn Ser Val Ala Val Ser Pro Asn Ala Ala Ile Ile Ser Thr Ala Ala Val Pro Gly Asp Ala Gly Ser Val Ala Ala Met Phe Gln Met Ser Pro Pro Leu Gln Ser Ala Pro Ser Ser Gln Asp Gly Val Ser Pro Gly Ser Gly Ser Asn Ser Ser Ser Ser Val Gly Val Phe Ser Val Gly Ser Gly Ser Ser Gly Gly Val Gly Val Ser Asn Asp Asn His Gly Ala Gly Gly Thr Ala Ala Val Ser Tyr Gln Gly Ala Thr Val Phe Glu Pro Glu Val Gly Tyr Tyr Asn Asp Pro Val Ala Val Phe Asp Phe Ala Ser Leu Tyr Pro Ser Ile Ile Met Ala His Asn Leu Cys Tyr Ser Thr Leu

Leu Val Pro Gly Gly Glu Tyr Pro Val Asp Pro Ala Asp Val Tyr Ser 740 745 750

- Val Thr Leu Glu Asn Gly Val Thr His Arg Phe Val Arg Ala Ser Val
 755 760 765
- Arg Val Ser Val Leu Ser Glu Leu Leu Asn Lys Trp Val Ser Gln Arg
 770 780
- Arg Ala Val Arg Glu Cys Met Arg Glu Cys Gln Asp Pro Val Arg Arg 785 790 795 800
- Met Leu Leu Asp Lys Glu Gln Met Ala Leu Lys Val Thr Cys Asn Ala 805 810 815
- Phe Tyr Gly Phe Thr Gly Ala Leu Asn Gly Met Met Pro Cys Leu Pro 820 825 830
- Ile Ala Ala Ser Ile Thr Arg Ile Gly Arg Asp Met Leu Glu Arg Thr 835 840 845
- Ala Arg Phe Ile Lys Asp Asn Phe Ser Glu Pro Cys Phe Leu His Asn 850 855 860
- Phe Phe Asn Gln Glu Asp Tyr Val Val Gly Thr Arg Glu Gly Asp Ser 865 870 875 880
- Glu Glu Ser Ser Ala Leu Pro Glu Gly Leu Glu Thr Ser Ser Gly Gly 895
- Ser Asn Glu Arg Arg Val Glu Ala Arg Val Ile Tyr Gly Asp Thr Asp 900 905 910
- Ser Val Phe Val Arg Phe Arg Gly Leu Thr Pro Gln Ala Leu Val Ala 915 920 925
- Arg Gly Pro Ser Leu Ala His Tyr Val Thr Ala Cys Leu Phe Val Glu 930 935 940
- Pro Val Lys Leu Glu Phe Glu Lys Val Phe Val Ser Leu Met Met Ile 945 950 955 960
- Cys Lys Lys Arg Tyr Ile Gly Lys Val Glu Gly Ala Ser Gly Leu Ser 975
- Met Lys Gly Val Asp Leu Val Arg Lys Thr Ala Cys Glu Phe Val Lys 980 985 990
- Gly Val Thr Arg Asp Val Leu Ser Leu Leu Phe Glu Asp Arg Glu Val 995 1000 1005
- Ser Glu Ala Ala Val Arg Leu Ser Arg Leu Ser Leu Asp Glu Val 1010 1015 1020
- Lys Lys Tyr Gly Val Pro Arg Gly Phe Trp Arg Ile Leu Arg Arg 1025 1030 1035
- Leu Val Gln Ala Arg Asp Asp Leu Tyr Leu His Arg Val Arg Val 1040 1045 1050
- Glu Asp Leu Val Leu Ser Ser Val Leu Ser Lys Asp Ile Ser Leu 1055 1060 1065

- Tyr Val Leu Thr Ala Pro Gly Val Arg Thr Ala Pro Gln Gly Ser 1100 1105 1110
- Ser Asp Asn Gly Asp Ser Val Thr Ala Gly Val Val Ser Arg Ser 1115 1120 1125
- Asp Ala Ile Asp Gly Thr Asp Asp Asp Ala Asp Gly Gly Gly Val 1130 1135 1140
- Glu Glu Ser Asn Arg Arg Gly Glu Pro Ala Lys Lys Arg Ala 1145 1150 1155
- Arg Lys Pro Pro Ser Ala Val Cys Asn Tyr Glu Val Ala Glu Asp 1160 1165 1170
- Pro Ser Tyr Val Arg Glu His Gly Val Pro Ile His Ala Asp Lys 1175 1180 1185
- Tyr Phe Glu Gln Val Leu Lys Ala Val Thr Asn Val Leu Ser Pro 1190 1195 1200
- Val Phe Pro Gly Gly Glu Thr Ala Arg Lys Asp Lys Phe Leu His 1205 1210 1215
- Met Val Leu Pro Arg Arg Leu His Leu Glu Pro Ala Phe Leu Pro 1220 1225 1230
- Tyr Ser Val Lys Ala His Glu Cys Cys 1235 1240
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- <211> 1242
- <212> PRT
- <213> herpes simplex
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- Ala Gly Gly Arg Arg Gln Arg Ser Gln Pro Gly Ser Ala Gln Gly Ser 20 25 30
- Gly Lys Arg Pro Pro Gln Lys Gln Phe Leu Gln Ile Val Pro Arg Gly 35 40 45
- Val Met Phe Asp Gly Gln Thr Gly Leu Ile Lys His Lys Thr Gly Arg
 50 55 60
- Leu Pro Leu Met Phe Tyr Arg Glu Ile Lys His Leu Leu Ser His Asp
 65 70 75 80
- Met Val Trp Pro Cys Pro Trp Arg Glu Thr Leu Val Gly Arg Val Val 85 90 95

Gly	Pro	Ile	Arg 100	Phe	His	Thr	Tyr	Asp 105	Gln	Thr	Asp	Ala	Val 110	Leu	Phe
Phe	Asp	Ser 115	Pro	Glu	Asn	Val	Ser 120	Pro	Arg	Tyr	Arg	Gln 125	His	Leu	Val
Pro	Ser 130	Gly	Asn	Val	Leu	Arg 135	Phe	Phe	Gly	Ala	Thr 140	Glu	His	Gly	Туг
Ser 145	Ile	Cys	Val	Asn	Val 150	Phe	Gly	Gln	Arg	Ser 155	Tyr	Phe	Tyr	Суѕ	Glu 160
Tyr	Ser	Asp	Thr	Asp 165	Arg	Leu	Arg	Glu	Val 170	Ile	Ala	Ser	Val	Gly 175	Glu
Leu	Val	Pro	Glu 180	Pro	Arg	Thr	Pro	Tyr 185	Ala	Val	Ser	Val	Thr 190	Pro	Ala
Thr	Lys	Thr 195	Ser	Ile	Tyr	Gly	Tyr 200	Gly	Thr	Arg	Pro	Val 205	Pro	Asp	Leu
Gln	**		Ser				Trp				Arg 220	Lys	Ile	Gly	Glu
Tyr 225	Leu	Leu	Glu	Gln	Gly 230	Phe	Pro	Val	Tyr	Glu 235	Val	Arg	Val	Asp	Pro 240
Leu	Thr	Arg	Leu	Val 245	Ile	Asp	Arg	Arg	Ile 250	Thr	Thr	Phe	Gly	Trp 255	Суѕ
Ser	Val	Asn	Arg 260	Tyr	Asp	Trp	Arg	Gln 265	Gln	Gly	Arg	Ala	Ser 270	Thr	Cys
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	290					295					300		Ile		
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			340					345					Cys 350		
		355					360					365	Thr		
	370					375					380		Phe		
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				405					410				Leu	415	
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Cys	Lys	Leu 435	Pro	Thr	Ala	Gln	Gly 440	Gly	Arg	Phe	Phe	Leu 445	His	Ser	Pro
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Leu 545		_			Phe 550										
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Arg	Asp	Phe 595	Ile	Leu	Pro	Asn	His 600	Tyr	Ser	Lys	Gly	Thr 605	Thr	Val	Pro
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Ala 625	Ala	Val	Pro	Gly	Asp 630	Ala	Gly	Ser	Val	Ala 635	Ala	Met	Phe	Gln	Met
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Leu	Val	Pro	Gly 740	Gly	Glu	Tyr	Pro	Val 745	Asp	Pro	Ala	Asp	Val 750	Tyr	Ser
Va1	Thr	Len	Glu	Asn	Gly	Val	Thr	His	Ara	Phe	Val	Ara	Ala	Ser	Val

**	Q 0 _ .														
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Met	Leu	Leu	Asp	Lys 805	Glu	Gln	Met	Ala	Leu 810	Lys	Val	Thr	Cys	Asn 815	Ala
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Phe 865	Phe	Asn	Gln	Glu	Asp 870	Tyr	Val	Val	Gly	Thr 875	Arg	Glu	Gly	Asp	Ser 880
Glu	Glu	Ser	Ser	Ala 885	Leu	Pro	Glu	Gly	Leu 890	Glu	Thr	Ser	Ser	Gly 895	Gl _Y
,Ser	Asn	Glu	Arg 900	Arg	Val	Glu	Ala	Arg 905	Val	Ile	Tyr	Gly	Asp 910	Thr	Asp
Ser	Val	Phe 915	Val	Arg	Phe	Arg	Gly 920	Leu	Thr	Pro	Gln	Ala 925		Val	Ala
Arg	Gly 930	Pro	Ser	Leu	Ala	His 935	Tyr	Val	Thr	Ala	Cys 940	Leu	Phe	Val	Glu
Pro 945	Val	Lys	Leu	Glu	Phe 950	Glu	Ĺys	Val	Phe	Val 955	Ser	Leu	Met	Met	Ile 960
Суѕ	Lys	Lys	Arg	Tyr 965	Ile	Gly	ГÀЗ	Val	Glu 970	Gly	Ala	Ser	Gly	Leu 975	Ser
Met	Lys	Gly	Val 980	Asp	Leu	Val	Arg	Lys 985	Thr	Ala	Cys	Glu	Phe 990	Val	Lys
Gly	Val	Thr 995	Arg	Asp	Val	Leu	Ser		ı Le	u Phe	e Gl	u As		rg G	lu Val
Ser	Glu- 1010		a Ala	a Val	l Arg	J Let 10:		er A	rg L	eu Se		eu 020	Asp (Glu '	Val
Lys	Lys 1025	_	r Gl	y Val	l Pro	10:		ly P	he T	rp A		le 035	Leu Z	Arg /	Arg
Leu	Val 1040		n Ala	a Arg	g Ası	Ası 10		eu T	yr L	eu H		rg 050	Val Z	Arg '	Val
Glu	Asp 105		u Va	l Lei	ı Sei	10		al L	eu S	er L	_	sp 065	Ile :	Ser :	Leu
Tyr	Arg 107		n Se	r Ası	n Lei	10°		is I	le A	la V		le 080	Lys i	Arg :	Leu

Ala Ala Arg Ser Glu Glu Leu Pro Ser Val Gly Asp Arg Val Phe 1085 1090 1095

- Tyr Val Leu Thr Ala Pro Gly Val Arg Thr Ala Pro Gln Gly Ser 1100 1105 1110
- Ser Asp Asn Gly Asp Ser Val Thr Ala Gly Val Val Ser Arg Ser 1115 1120 1125
- Asp Ala Ile Asp Gly Thr Asp Asp Asp Ala Asp Gly Gly Val 1130 1135 1140
- Glu Glu Ser Asn Arg Arg Gly Gly Glu Pro Ala Lys Lys Arg Ala 1145 1150 1155
- Arg Lys Pro Pro Ser Ala Val Cys Asn Tyr Glu Val Ala Glu Asp 1160 1165 1170
- Pro Ser Tyr Val Arg Glu His Gly Val Pro Ile His Ala Asp Lys 1175 1180 1185
- Tyr Phe Glu Gln Val Leu Lys Ala Val Thr Asn Val Leu Ser Pro 1190 1195 1200
- Val Phe Pro Gly Gly Glu Thr Ala Arg Lys Asp Lys Phe Leu His 1205 1210 1215
- Met Val Leu Pro Arg Arg Leu His Leu Glu Pro Ala Phe Leu Pro 1220 1230
- Tyr Ser Val Lys Ala His Glu Cys Cys 1235 1240
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- Ala Arg Ala Ala Ser Gly Phe Phe Ala Pro His Asn Pro Arg Gly Ala 20 25 30
- Thr Gln Thr Ala Pro Pro Pro Cys Arg Arg Gln Asn Phe Tyr Asn Pro 35 40 45
- His Leu Ala Gln Thr Gly Thr Gln Pro Lys Ala Pro Gly Pro Ala Gln 50 55 60
- Arg His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro 65 70 . 75 80
- Arg Ser Leu Asp Glu Asp Ala Pro Ala Glu Gln Arg Thr Gly Val His
 85 90 95
- Asp Gly Arg Leu Arg Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu 100 105 110
- Arg Asp Val Leu Arg Val Gly Pro Glu Gly Phe Trp Pro Arg Arg Leu

		115					120					125			
Arg	Leu 130	Trp	Gly	Gly	Ala	Asp 135	His	Ala	Pro	Lys	Gly 140	Phe	Asp	Pro	Thr
Val 145	Thr	Val	Phe	His	Val 150	Tyr	Asp	Ile	Leu	Glu 155	His	Val	Glu	His	Ala 160
Tyr	Ser	Met	Arg	Ala 165	Ala	Gln	Leu	His	Glu 170	Arg	Phe	Met	Asp	Ala 175	Ile
Thr	Pro	Ala	Gly 180	Thr	Val	Ile	Thr	Leu 185	Leu	Gly	Leu	Thr	Pro 190	Glu	Gly
His	Arg	Val 195	Ala	Val	His	Val	Tyr 200	Gly	Thr	Arg	Gln	Tyr 205	Phe	Tyr	Met
Ásn	Lys 210	Ala	Glu	Val	Asp	Arg 215	His	Leu	Glń	Cys	Arg 220	Ala	Pro	Arg	Asp
Leu 225	Cys	Glu	Arg	Leu	Ala 230	Ala	Ala	Leu	Arg	Glu 235	Ser	Pro	Gly	Ala	Ser 240
Phe	Arg	Gly	Ile	Ser 245	Ala	Asp	His	Phe	Glu 250	Ala	Glu	Val	Val	Glu 255	Arg
Ala	Asp	Val	Tyr 260	Tyr	Tyr	Glu	Thr	Arg 265	Pro	Thr	Leu	Tyr	Tyr 270	Arg	Val
Phe	Val	Arg 275	Ser	Gly	Arg	Ala	Leu 280	Ala	Tyr	Leu	Cys	Asp 285	Asn	Phe	Cys
Pro	Ala 290	Ile	Arg	Lys	Tyr	Glu 295	Gly	Gly	Val	Asp	Ala 300	Thr	Thr	Arg	Phe
Ile 305	Leu	Asp	Asn	Pro	Gly 310	Phe	Val	Thr	Phe	Gly 315	Trp	Tyr	Arg	Leu	Lys 320
Pro	Gly	Arg	Gly	Asn 325	Ala	Pro	Ala	Gln	Pro 330	Arg	Pro	Pro	Thr	Ala 335	Phe
Gly	Thr	Ser	Ser 340	Asp	Val	Glu	Phe	Asn 345	Cys	Thr	Ala	Asp	Asn 350	Leu	Ala
Val	Glu	Gly 355	Ala	Met	Cys	Asp	Leu 360	Pro	Ala	Tyr	Lys	Leu 365	Met	Cys	Phe
Asp	Ile 370	Glu	Cys	Lys	Ala	Gly 375	Gly	Glu	Asp	Glu	Leu 380	Ala	Phe	Pro	Val
Ala 385	Glu	Arg	Pro	Glu	Asp 390	Leu	V al	Ile	Gln	Ile 395	Ser	Cys	Leu	Leu	Tyr 400
Asp	Leu	Ser	Thr	Thr 405	Ala	Leu	Glu	His	Ile 410	Leu	Leu	Phe	Ser	Leu 415	Gly
Ser	Суѕ	Asp	Leu 420	Pro	Glu	Ser	His	Leu 425	Ser	Asp	Leu	Ala	Ser 430	Arg	Gly
Leu	Pro	Ala	Pro	Val	Val	Leu	Glu	Phe	Asp	Ser	Glu	Phe	Glu	Met	Leu

Leu	Ala 450	Phe	Met	Thr	Phe	Val 455	Lys	Gln	Tyr	Gly	Pro 460	Glu	Phe	Val	Thr
Gly 465	Tyr	Asn	Ile	Ile	Asn 470	Phe	Asp	Trp	Pro	Phe 475	Val	Leu	Thr	Lys	Leu 480
Thr	Glu	Ile	Tyr	Lys 485	Val	Pro	Leu	Asp	Gly 490	Tyr	Gly	Arg	Met	Asn 495	Gly
Arg	Gly	Val	Phe 500	Arg	Val	Trp	Asp	Ile 505	Gly	Gln	Ser	His	Phe 510	Gln	Lys
Arg	Ser	Lys 515	Ile	Lys	Val	Asn	Gly 520	Met	Val	Asn	Ile	Asp 525	Met	Tyr	Gly
Ile	Ile 530	Thr	Asp	Lys	Val	Lys 535	Leu	Ser	Ser	Tyr	Lys 540	Leu	Asn	Ala	Val
Ala 545	Glu	Ala	Val	Leu	Lys 550	Asp	Lys	Lys	Lys	Asp 555	Leu	Ser	Tyr	Arg	Asp 560
Ile	Pro	Ala	Tyr		Ala			Pro		Gln				Ile 575	Gly
Glu	Tyr	Cys	Val 580	Gln	Asp	Ser	Leu	Leu 585	Val	Gly	Gln	Leu	Phe 590	Phe	Lys
Phe	Leu	Pro 595	His	Leu	Glu	Leu	Ser 600	Ala	Val	Ala	Arg	Leu 605	Ala	Gly	Ile
Asn	Ile 610	Thr	Arg	Thr	Ile	Tyr 615	Asp	Gly	Gln	Gln	Ile 620	Arg	Val	Phe	Thr
Cys 625	Leu	Leu	Arg	Leu	Ala 630	Gly	Gln	Lys	Gly	Phe 635	Ile	Leu	Pro	Asp	Thr 640
Gln	Gly	Arg	Phe	Arg 645	Gly	Leu	Asp	ГЛЗ	Glu 650	Ala	Pro	Lys	Arg	Pro 655	Ala
Val	Pro	Arg	Gly 660	Glu	Gly	Glu	Arg	Pro 665	Gly	Asp	Gly	Asn	Gly 670	Asp	Glu
Asp	Lys	Asp 675	Asp	Asp	Glu	Asp	Glu 680	Asp	Gly	Asp	Glu	Arg 685	Glu	Glu	Val
Ala	Arg 690	Glu	Thr	Gly	Gly	Arg 695	His	Val	Gly	Tyr	Gln 700	Gly	Ala	Arg	Val
Leu 705	Asp	Pro	Thr	Ser	Gly 710	Phe	His	Val	Asp	Pro 715	Val	Val	Val	Phe	Asp 720
Phe	Ala	Ser	Leu	Tyr 725	Pro	Ser	Ile	Ile	.Gln 730	Ala	His	Asn	Leu	Cys 735	Phe
Ser	Thr	Leu	Ser 740	Leu	Arg	Pro	Glu	Ala 745	Val	Ala	His	Leu	Glu 750	Ala	Asp
Arg	Asp	Tyr 755	Leu	Glu	Ile	Glu	Val 760	Gly	Gly	Arg	Arg	Leu 765	Phe	Phe	Val
Lys	Ala 770	His	Val	Arg	Glu	Ser 775	Leu	Leu	Ser	Ile	Leu 780	Leu	Arg	Asp	Trp

Leu Ala Met Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Thr Pro 785 790 795 800

- Glu Glu Ala Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val 805 810 815
- Cys Asn Ser Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu Leu Pro 820 825 830
- Cys Leu His Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu . 835 840 845
- Leu Ala Thr Arg Ala Tyr Val His Ala Arg Trp Ala Glu Phe Asp Gln 850 855 860
- Leu Leu Ala Asp Phe Pro Glu Ala Ala Gly Met Arg Ala Pro Gly Pro 875 880
- Tyr Ser Met Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu 885 890 895
- Cys Arg Gly Leu Thr Ala Ala Gly Leu Val Ala Met Gly Asp Lys Met 900 905 . 910
- Ala Ser His Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu 915 920 925
- Cys Glu Lys Thr Phe Thr Lys Leu Leu Ile Ala Lys Lys Lys Tyr 930 935 940
- Ile Gly Val Ile Cys Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu 945 950 955 960
- Val Arg Lys Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu 965 970 975
- Val Asp Leu Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala 980 985 990
- Leu Ala Glu Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu 995 1000 1005
- Gly Leu Gln Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg 1010 1015 1020
- Ile Thr Asp Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala 1025 1030 1035
- Glu Leu Ser Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala 1040 1045 1050
- His Leu Thr Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val 1055 1060 1065
- Pro Ser Ile Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr 1070 1075 1080
- Arg Glu Val Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu 1085 1090 1095
- Leu Asp Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala

: ' .::

PCT/US01/16525

WO 02/06513 Leu Pro Ser Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser His Ala Asp Pro Pro Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu Val Ser Glu Leu Ala Glu Asp Pro Gly Tyr Ala Ile Ala Arg Gly Val Pro Leu Asn Thr Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala Ala Cys Val Thr Phe Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile Thr Glu Ser Leu Leu Lys Arg Phe Ile Pro Glu Thr Trp His Pro Pro Asp Asp Val Ala Ala Arg Leu Arg Ala Ala Gly Phe Gly Pro Ala Gly Ala Gly Ala Thr Ala Glu Glu Thr Arg Arg Met Leu His Arg Ala Phe Asp Thr Leu Ala <210> 15 <211> 1240 <212> PRT <213> herpes simplex <400> 15 Met Phe Cys Ala Ala Gly Gly Pro Ala Ser Pro Gly Gly Lys Ser Ala Ala Arg Ala Ala Ser Gly Phe Phe Ala Pro His Asn Pro Arg Gly Ala Thr Gln Thr Ala Pro Pro Pro Cys Arg Arg Gln Asn Phe Tyr Asn Pro His Leu Ala Gln Thr Gly Thr Gln Pro Lys Ala Pro Gly Pro Ala Gln Arg His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg Ser Leu Asp Glu Asp Ala Pro Ala Glu Gln Arg Thr Gly Val His

Asp Gly Arg Leu Arg Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu Arg Asp Val Leu Arg Val Gly Pro Glu Gly Phe Trp Pro Arg Arg Leu Arg Leu Trp Gly Gly Ala Asp His Ala Pro Glu Gly Phe Asp Pro Thr

Val 145	Thr	Val	Phe	His	Val 150	Tyr	Asp	Ile	Leu	Glu 155	His	Val	Glu	His	Ala 160
Tyr	Ser	Met	Arg	Ala 165	Ala	Gln	Leu	His	Glu 170	Arg	Phe	Met	Asp	Ala 175	Ile
Thr	Pro	Ala	Gly 180	Thr	Val	Ile	Thr	Leu 185	Leu	Gly	Leu	Thr	Pro 190	Glu	Gly
His	Arg	Val 195	Ala	Val	His	Val	Tyr 200	Gly	Thr	Arg	Gln	Tyr 205	Phe	Tyr	Met
Asn	Lys 210	Ala	Glu	Val	Asp	Arg 215	His	Leu	Gln	Cys	Arg 220	Ala	Pro	Arg	Asp
Leu 225	Cys	Glu	Arg	Leu	Ala 230	Ala	Ala	Leu	Arg	Glu 235	Ser	Pro	Gly	Ala	Ser 240
Phe	Arg	Gly	Ile	Ser 245	Ala	Asp	His	Phe	Glu 250	Ala	Glu	Val	Val	Glu 255	Arg
Ala	Asp	Val			Tyr								Tyr 270	Arg	Val
Phe	Val	Arg 275	Ser	Gly	Arg	Ala	Leu 280	Ala	Tyr	Leu	Суѕ	Asp 285	Asn	Phe	Cys
Pro	Ala 290	Ile	Arg	Lys	Tyr	Glu 295	Gly	Gly	Val	Asp	Ala 300	Thr	Thr	Arg	Phe
Ile 305	Leu	Asp	Asn	Pro	Gly 310	Phe	Val	Thr	Phe	Gly 315	Trp	Tyr	Arg	Leu	Lys 320
Pro	Gly	Arg	Gly	Asn 325	Ala	Pro	Ala	Gln	Pro 330	Arg	Pro	Pro	Thr	Ala 335	Phe
Gly	Thr	Ser	Ser 340	Asp	Val	Glu	Phe	Asn 345	Cys	Thr	Ala	Asp	Asn 350	Leu	Ala
Val	Glu	Gly 355	Ala	Met	Cys	Asp	Leu 360	Pro	Ala	Tyr	Lys	Leu 365	Met	Cys	Phe
Asp	Ile 370	Glu	Cys	Lys	Ala	Gly 375	Gly	Glu	Asp	Glu	Leu 380	Ala	Phe	Pro	Val
Ala 385	Glu	Arg	Pro	Glu	Asp 390	Leu	Val	Ile	Gln	Ile 395	Ser	Cys	Leu	Leu	Tyr 400
Asp	Leu	Ser	Thr	Thr 405	Ala	Leu	Glu	His	Ile 410	Leu	Leu	Phe	Ser	Leu 415	Gly
Ser	Cys	Asp	Leu 420	Pro	Glu	Ser	His	Leu 425	Ser	Asp	Leu	Ala	Ser 430	Arg	Gly
Leu	Pro	Ala 435	Pro	Val	Val	Leu	Glu 440	Phe	Asp	Ser	Glu	Phe 445	Glu	Met	Leu
Leu	Ala 450	Phe	Met	Thr	Phe	Val 455	Lys	Gln	Tyr	Gly	Pro 460	Glu	Phe	Val	Thr
Clar	Пчет	λαν	Tla	T10	Δsn	Dhe	ħ cr	m-r-r	Dro	Dhe	17a 1	T 011	mb~	Laze	T.em

465					470					475					480
Thr	Glu	Ile	Tyr	Lys 485	Val	Pro	Leu	Asp	Gly 490	Tyr	Gly	Arg	Met	Asn 495	Gly
Arg	Gly	Val	Phe 500	Arg	Val	Trp	Asp	Ile 505	Gly	Gln	Ser	His	Phe 510	Gln	Lys
Arg	Ser	Lys 515	Ile	Lys	Val	Asn	Gly 520	Met	Val	Asn	Ile	Asp 525	Met	Tyr	Gly
Ile	Ile 530	Thr	Asp	Lys	Val	Lys 535	Leu	Ser	Ser	Tyr	Lys 540	Leu	Asn	Ala	Val
Ala 545	Glu	Ala	Val	Leu	Lys 550	Asp	Lys	Lys	Lys	Asp 555	Leu	Ser	Tyr	Arg	Asp 560
Iļē	Pro	Ala	Tyr	Tyr 565	Ala	Ser	Gly	Pro	Ala 570	Gln	Arg	Gly	Val	Ile 575	
Glu	Tyr	Суз	Val 580	Gln	Asp	Ser	Leu	Leu 585	Val	Gly	Gln	Leu	Phe 590	Phe	Lys
Phe	Leu	Pro 595	His	Leu	Glu	Leu	Ser 600	Ala	Val	Ala	Arg	Leu 605	Ala	Gly	Ile
Asn	Ile 610	Thr	Arg	Thr	Ile	Tyr 615	Asp	Gly	Gln	Gln	Ile 620	Arg	Val	Phe	Thr
Cys 625	Leu	Leu	Arg	Leu	Ala 630	Gly	Gln	Lys	Gly	Phe 635	Ile	Leu	Pro	qzA	Thr 640
Gln	Gly	Arg	Phe	Arg 645	Gly	Leu	Asp	Lys	Glu 650	Ala	Pro	Lys	Arg	Pro 655	Ala
Val	Pro	Arg	Gly 660	Glu	Gly	Glu	Arg	Pro 665	Gly	Asp	Gly	Asn	Gly 670	Asp	Glu
Asp	Lys	Asp 675	Asp	Asp	Glu	Asp	Gly 680	Asp	Glu	Asp	Gly	Asp 685	Glu	Arg	Glu
Glu	Val 690	Ala	Arg	Glu	Thr	Gly 695	Gly	Arg	His	Val	Gly 700	Tyr	Gln	Gly	Ala
Arg 705	Val	Leu	Asp	Pro	Thr 710	Ser	Gly	Phe	His	Val 715	Asp	Pro	Val	Val	Val 720
Phe	Asp	Phe	Ala	Ser 725	Leu	Tyr	Pro	Ser	Ile 730	Ile	Gln	Ala	His	Asn 735	Leu
Cys	Phe	Ser	Thr 740	Leu	Ser	Leu	Arg	Pro 745	Glu	Ala	Val	Ala	His 750	Leu	Glu
Ala	Asp	Arg 755	Asp	Tyr	Leu	Glu	Ile 760	Glu	Val	Gly	Gly	Arg 765	Arg	Leu	Phe
Phe	Val 770	Lys	Ala	His	Val	Arg 775	Glu	Ser	Leu	Leu	Ser 780	Ile	Leu	Leu	Arg
Asp 785	Trp	Leu	Ala	Met	Arg 7.90	Lys	Gln	Ile	Arg	Ser 795	Arg	Ile	Pro	Gln	Ser 800

Pro Pro Glu Glu Ala Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys 805 810 815

- Val Val Cys Asn Ser Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu 820 825 830
- Leu Pro Cys Leu His Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu 835 840 845
- Met Leu Leu Ala Thr Arg Ala Tyr Val His Ala Arg Trp Ala Glu Phe 850 855 860
- Asp Gln Leu Leu Ala Asp Phe Pro Glu Ala Ala Gly Met Arg Ala Pro 865 870 875 880
- Gly Pro Tyr Ser Met Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe 885 890 895
- Val Leu Cys Arg Gly Leu Thr Ala Ala Gly Leu Val Ala Met Gly Asp 900 905 910
- Lys Met Ala Ser His Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys 915 920 925
- Leu Glu Cys Glu Lys Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys 930 935 940
- Lys Tyr Ile Gly Val Ile Cys Gly Gly Lys Met Leu Ile Lys Gly Val 945 950 955 960
- Asp Leu Val Arg Lys Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg 965 970 975
- Ala Leu Val Asp Leu Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala 980 985 990
- Ala Ala Leu Ala Glu Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu 995 1000 1005
- Pro Glu Gly Leu Gln Ala Phe Gly Ala Val Leu Val Asp Ala His 1010 1015 1020
- Arg Arg Ile Thr Asp Pro Glu Arg Asp Ile Gln Asp Phe Val Leu 1025 1030 1035
- Thr Ala Glu Leu Ser Arg His Pro Arg Ala Tyr Thr Asn Lys Arg 1040 1045 1050
- Leu Ala His Leu Thr Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala 1055 1060 1065
- Gln Val Pro Ser Ile Lys Asp Arg Ile Pro Tyr Val Ile Val Ala 1070 1075 1080
- Gln Thr Arg Glu Val Glu Glu Thr Val Ala Arg Leu Ala Ala Leu 1085 1090 1095
- Arg Glu Leu Asp Ala Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro
 1100 1105 1110
- Ala Ala Leu Pro Ser Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser 1115 1120 1125

His Ala Asp Pro Pro Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu 1130 1135 1140

- Val Ser Glu Leu Ala Glu Asp Pro Gly Tyr Ala Ile Ala Arg Gly 1145 1150 1155
- Val Pro Leu Asn Thr Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala 1160 1165 1170
- Ala Cys Val Thr Phe Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile 1175 1180 1185
- Thr Glu Ser Leu Leu Lys Arg Phe Ile Pro Glu Thr Trp His Pro 1190 1195 1200
- Pro Asp Asp Val Ala Ala Arg Leu Arg Ala Ala Gly Phe Gly Pro 1205: 1215
- Ala Gly Ala Gly Ala Thr Ala Glu Glu Thr Arg Arg Met Leu His 1220 1225 1230
- Arg Ala Phe Asp Thr Leu Ala 1235 1240
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- Gly Arg Gly Pro Pro Pro Cys Leu Arg Gln Asn Phe Tyr Asn Pro Tyr 35 40 . 45
- Leu Ala Pro Val Gly Thr Gln Gln Lys Pro Thr Gly Pro Thr Gln Arg 50 55 60
- His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 75 80
- Val Leu Asp Glu Asp Ala Pro Pro Glu Lys Arg Ala Gly Val His Asp 85 90 95
- Gly His Leu Lys Arg Ala Pro Lys Val. Tyr Cys Gly Gly Asp Glu Arg
 100 105 110
- Asp Val Leu Arg Val Gly Ser Gly Gly Phe Trp Pro Arg Arg Ser Arg 115 120 125
- Leu Trp Gly Gly Val Asp His Ala Pro Ala Gly Phe Asn Pro Thr Val
- Thr Val Phe His Val Tyr Asp Ile Leu Glu Asn Val Glu His Ala Tyr 145 150 150

Gly Met Arg Ala Ala Gln Phe His Ala Arg Phe Met Asp Ala Ile Thr Pro Thr Gly Thr Val Ile Thr Leu Leu Gly Leu Thr Pro Glu Gly His Arg Val Ala Val His Val Tyr Gly Thr Arg Gln Tyr Phe Tyr Met Asn Lys Glu Glu Val Asp Arg His Leu Gln Cys Arg Ala Pro Arg Asp Leu Cys Glu Arg Met Ala Ala Ala Leu Arg Glu Ser Pro Gly Ala Ser Phe Arg Gly Ile Ser Ala Asp His Phe Glu Ala Glu Val Val Glu Arg Thr Asp Val Tyr Tyr Tyr Glu Thr Arg Pro Ala Leu Phe Tyr Arg Val Tyr Val Arg Ser Gly Arg Val Leu Ser Tyr Leu Cys Asp Asn Phe Cys Pro Ala Ile Lys Lys Tyr Glu Gly Gly Val Asp Ala Thr Thr Arg Phe Ile Leu Asp Asn Pro Gly Phe Val Thr Phe Gly Trp Tyr Arg Leu Lys Pro Gly Arg Asn Asn Thr Leu Ala Gln Pro Arg Ala Pro Met Ala Phe Gly Thr Ser Ser Asp Val Glu Phe Asn Cys Thr Ala Asp Asn Leu Ala Ile Glu Gly Gly Met Ser Asp Leu Pro Ala Tyr Lys Leu Met Cys Phe Asp Ile Glu Cys Lys Ala Gly Gly Glu Asp Glu Leu Ala Phe Pro Val Ala Gly His Pro Glu Asp Leu Val Ile Gln Ile Ser Cys Leu Leu Tyr Asp Leu Ser Thr Thr Ala Leu Glu His Val Leu Leu Phe Ser Leu Gly Ser Cys Asp Leu Pro Glu Ser His Leu Asn Glu Leu Ala Ala Arg Gly Leu Pro Thr Pro Val Val Leu Glu Phe Asp Ser Glu Phe Glu Met Leu Leu Ala Phe Met Thr Leu Val Lys Gln Tyr Gly Pro Glu Phe Val Thr Gly Tyr Asn Ile Ile Asn Phe Asp Trp Pro Phe Leu Leu Ala Lys Leu Thr Asp Ile Tyr Lys Val Pro Leu Asp Gly Tyr Gly Arg Met Asn Gly Arg

Gly Val Phe Arg Val Trp Asp Ile Gly Gln Ser His Phe Gln Lys Arg Ser Lys Ile Lys Val Asn Gly Met Val Asn Ile Asp Met Tyr Gly Ile Ile Thr Asp Lys Ile Lys Leu Ser Ser Tyr Lys Leu Asn Ala Val Ala Glu Ala Val Leu Lys Asp Lys Lys Asp Leu Ser Tyr Arg Asp Ile Pro Ala Tyr Tyr Ala Ala Gly Pro Ala Gln Arg Gly Val Ile Gly Glu Tyr Cys Ile Gln Asp Ser Leu Leu Val Gly Gln Leu Phe Phe Lys Phe 59Û Leu Pro His Leu Glu Leu Ser Ala Val Ala Arg Leu Ala Gly Ile Asn Ile Thr Arg Thr Ile Tyr Asp Gly Gln Gln Ile Arg Val Phe Thr Cys Leu Leu Arg Leu Ala Asp Gln Lys Gly Phe Ile Leu Pro Asp Thr Gln Gly Arg Phe Arg Gly Ala Gly Glu Ala Pro Lys Arg Pro Ala Ala Ala Arg Glu Asp Glu Glu Arg Pro Glu Glu Glu Gly Glu Asp Glu Asp Glu Arg Glu Glu Gly Gly Glu Arg Glu Pro Glu Gly Ala Arg Glu Thr Ala Gly Arg His Val Gly Tyr Gln Gly Ala Arg Val Leu Asp Pro Thr Ser Gly Phe His Val Asn Pro Val Val Val Phe Asp Phe Ala Ser Leu Tyr Pro Ser Ile Ile Gln Ala His Asn Leu Cys Phe Ser Thr Leu Ser Leu Arg Ala Asp Ala Val Ala His Leu Glu Ala Gly Lys Asp Tyr Leu Glu Ile Glu Val Gly Gly Arg Arg Leu Phe Phe Val Lys Ala His Val Arg Glu Ser Leu Leu Ser Ile Leu Leu Arg Asp Trp Leu Ala Met Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val Cys Asn Ser Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu Leu Pro Cys Leu His

820 825 830

Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu Leu Ala Thr 835 840 845

- Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala 850 860
- Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met 865 870 875 880
- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 885 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Met Gly Asp Lys Met Ala Ser His
 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys945950950955
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr
 1040 1050
- Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala 1085 1090 1095
- Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala Leu Pro Ser 1100 1105 1110
- Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser His Ala Asp Pro Pro 1115 1120 1125
- Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu Val Ser Glu Leu Ala 1130 1135 1140

Glu Asp Pro Ala Tyr Ala Ile Ala His Gly Val Ala Leu Asn Thr 1145 1150 1155

- Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala Ala Cys Val Thr Phe 1160 1165 1170
- Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile Thr Glu Ser Leu Leu 1175 1180 1185
- Lys Arg Phe Ile Pro Glu Val Trp His Pro Pro Asp Asp Val Ala 1190 1195 1200
- Ala Arg Leu Arg Ala Ala Gly Phe Gly Ala Val Gly Ala Gly Ala 1205 1210 1215
- Thr Ala Glu Glu Thr Arg Arg Met Leu His Arg Ala Phe Asp Thr 1220 1225 1230

Leu Ala 1235

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<213> herpes simplex

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- Ala Arg Ala Ala Ser Gly Phe Phe Ala Pro Ala Gly Pro Arg Gly Ala 20 25 30
- Gly Arg Gly Pro Pro Pro Cys Leu Arg Gln Asn Phe Tyr Asn Pro Tyr 35 40 45
- Leu Ala Pro Val Gly Thr Gln Gln Lys Pro Thr Gly Pro Thr Gln Arg
 50 . 60
- His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 . 75 80
- Val Leu Asp Glu Asp Ala Pro Pro Glu Lys Arg Ala Gly Val His Asp 85 90 95
- Gly His Leu Lys Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu Arg
 100 105 110
- Asp Val Leu Arg Val Gly Ser Gly Gly Phe Trp Pro Arg Arg Ser Arg 115 120 125
- Leu Trp Gly Gly Val Asp His Ala Pro Ala Gly Phe Asn Pro Thr Val 130 135 140
- Thr Val Phe His Val Tyr Asp Ile Leu Glu Asn Val Glu His Ala Tyr 145 150 150
- Gly Met Arg Ala Ala Gln Phe His Ala Arg Phe Met Asp Ala Ile Thr 165 170 175
- Pro Thr Gly Thr Val Ile Thr Leu Leu Gly Leu Thr Pro Glu Gly His

190

Arg Val Ala Val His Val Tyr Gly Thr Arg Gln Tyr Phe Tyr Met Asn 195 200 205

185

180

Lys Glu Glu Val Asp Arg His Leu Gln Cys Arg Ala Pro Arg Asp Leu 210 220

Cys Glu Arg Met Ala Ala Ala Leu Arg Glu Ser Pro Gly Ala Ser Phe 225 230 235 240

Arg Gly Ile Ser Ala Asp His Phe Glu Ala Glu Val Val Glu Arg Thr 245 250 255

Asp Val Tyr Tyr Glu Thr Arg Pro Ala Leu Phe Tyr Arg Val Tyr 260 265 270

Val Arg Ser Gly Arg Val Leu Ser Tyr Leu Cys Asp Asm Phe Cys Pro 275 280 285

Ala Ile Lys Lys Tyr Glu Gly Gly Val Asp Ala Thr Thr Arg Phe Ile 290 295 300

Leu Asp Asn Pro Gly Phe Val Thr Phe Gly Trp Tyr Arg Leu Lys Pro 305 310 315 320

Gly Arg Asn Asn Thr Leu Ala Gln Pro Arg Ala Pro Met Ala Phe Gly 325 330 335

Thr Ser Ser Asp Val Glu Phe Asn Cys Thr Ala Asp Asn Leu Ala Ile 340 345 350

Glu Gly Gly Met Ser Asp Leu Pro Ala Tyr Lys Leu Met Cys Phe Asp 355 360 365

Ile Glu Cys Lys Ala Gly Gly Glu Asp Glu Leu Ala Phe Pro Val Ala 370 380

Gly His Pro Glu Asp Leu Val Ile Gln Ile Ser Cys Leu Leu Tyr Asp 385 390 395 400

Leu Ser Thr Thr Ala Leu Glu His Val Leu Leu Phe Ser Leu Gly Ser 405 410 415

Cys Asp Leu Pro Glu Ser His Leu Asn Glu Leu Ala Ala Arg Gly Leu
420 425 430

Pro Thr Pro Val Val Leu Glu Phe Asp Ser Glu Phe Glu Met Leu Leu 435 440 445

Ala Phe Met Thr Leu Val Lys Gln Tyr Gly Pro Glu Phe Val Thr Gly
450 460

Tyr Asn Ile Ile Asn Phe Asp Trp Pro Phe Leu Leu Ala Lys Leu Thr 465 470 475 480

Asp Ile Tyr Lys Val Pro Leu Asp Gly Tyr Gly Arg Met Asn Gly Arg
485
490
495

Gly Val Phe Arg Val Trp Asp Ile Gly Gln Ser His Phe Gln Lys Arg
500 505 510

Ser Lys Ile Lys Val Asn Gly Met Val Asn Ile Asp Met Tyr Gly Ile Ile Thr Asp Lys Ile Lys Leu Ser Ser Tyr Lys Leu Asn Ala Val Ala Glu Ala Val Leu Lys Asp Lys Lys Lys Asp Leu Ser Tyr Arg Asp Ile Pro Ala Tyr Tyr Ala Ala Gly Pro Ala Gln Arg Gly Val Ile Gly Glu Tyr Cys Ile Gln Asp Ser Leu Leu Val Gly Gln Leu Phe Phe Lys Phe Leu Pro His Leu Glu Leu Ser Ala Val Ala Arg Leu Ala Gly Ile Asn Ile Thr Arg Thr Ile Tyr Asp Gly Gln Gln Ile Arg Val Phe Thr Cys Leu Leu Arg Leu Ala Asp Gln Lys Gly Phe Ile Leu Pro Asp Thr Gln Gly Arg Phe Arg Gly Ala Gly Gly Glu Ala Pro Lys Arg Pro Ala Ala Ala Arg Glu Asp Glu Glu Arg Pro Glu Glu Glu Gly Glu Asp Glu Asp Glu Arg Glu Glu Gly Gly Gly Glu Arg Glu Pro Glu Gly Ala Arg Glu Thr Ala Gly Arg His Val Gly Tyr Gln Gly Ala Arg Val Leu Asp Pro Ile Ser Gly Phe His Val Asn Pro Val Val Val Phe Asp Phe Ala Ser Leu Tyr Pro Ser Ile Ile Gln Ala His Asn Leu Cys Phe Ser Thr Leu Ser Leu Arg Ala Asp Ala Val Ala His Leu Glu Ala Gly Lys Asp Tyr Leu Glu Ile Glu Val Gly Gly Arg Arg Leu Phe Phe Val Lys Ala His Val Arg Glu Ser Leu Leu Ser Ile Leu Leu Arg Asp Trp Leu Ala Met Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val Cys Asn Ser Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu Leu Pro Cys Leu His Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu Leu Ala Thr

Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala 850 855 860

- Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met 865 870 875 880
- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 885 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Met Gly Asp Lys Met Ala Ser His
 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys 945 950 955 960
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr 1040 1045 1050
- Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala 1085 1090 1095
- Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala Leu Pro Ser 1100 1105 1110
- Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser Pro Ala Asp Pro Pro 1115 1120 1125
- Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu Val Ser Glu Leu Ala 1130 1135 1140
- Glu Asp Pro Ala Tyr Ala Ile Ala His Gly Val Ala Leu Asn Thr 1145 1150 1155
- Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala Ala Cys Val Thr Phe

1160 1165 1170

Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile Thr Glu Ser Leu Leu 1175 1180 1185

Lys Arg Phe Ile Pro Glu Val Trp His Pro Pro Asp Asp Val Thr 1190 1195 1200

Ala Arg Leu Arg Ala Ala Gly Phe Gly Ala Val Gly Ala Gly Ala 1205 1210 1215

Thr Ala Glu Glu Thr Arg Arg Met Leu His Arg Ala Phe Asp Thr 1220 1230

Leu Ala 1235

<210> 18

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<212> PRT

<213> herpes simplex

<400> 18

Met Phe Ser Gly Gly Gly Pro Leu Ser Pro Gly Gly Lys Ser Ala
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Ala Arg Ala Ala Ser Gly Phe Phe Ala Pro Ala Gly Pro Arg Gly Ala 20 25 30

Gly Arg Gly Pro Pro Pro Cys Leu Arg Gln Asn Phe Tyr Asn Pro Tyr 35 40 45

Leu Ala Pro Val Gly Thr Gln Gln Lys Pro Thr Gly Pro Thr Gln Arg 50 55 60

His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 75 80

Val Leu Asp Glu Asp Ala Pro Pro Glu Lys Arg Ala Gly Val His Asp 85 90 95

Gly His Leu Lys Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu Arg 100 105 110

Asp Val Leu Arg Val Gly Ser Gly Gly Phe Trp Pro Arg Arg Ser Arg 115 120 125

Leu Trp Gly Gly Val Asp His Ala Pro Ala Gly Phe Asn Pro Thr Val 130 135 140

Thr Val Phe His Val Tyr Asp Ile Leu Glu Asn Val Glu His Ala Tyr 145 150 155 160

Gly Met Arg Ala Ala Gln Phe His Ala Arg Phe Met Asp Ala Ile Thr 165 170 175

Pro Thr Gly Thr Val Ile Thr Leu Leu Gly Leu Thr Pro Glu Gly His
180 185 190

Arg Val Ala Val His Val Tyr Gly Thr Arg Gln Tyr Phe Tyr Met Asn 195 200 205

Lys Glu Glu Val Asp Arg His Leu Gln Cys Arg Ala Pro Arg Asp Leu Cys Glu Arg Met Ala Ala Ala Leu Arg Glu Ser Pro Gly Ala Ser Phe Arg Gly Ile Ser Ala Asp His Phe Glu Ala Glu Val Val Glu Arg Thr Asp Val Tyr Tyr Tyr Glu Thr Arg Pro Ala Leu Phe Tyr Arg Val Tyr Val Arg Ser Gly Arg Val Leu Ser Tyr Leu Cys Asp Asn Phe Cys Pro Ala Ile Lys Lys Tyr Glu Gly Gly Val Asp Ala Thr Thr Arg Phe Ile Leu Asp Asn Pro Gly Phe Val Thr Phe Gly Trp Tyr Arg Leu Lys Pro Gly Arg Asn Asn Thr Leu Ala Gln Pro Arg Ala Pro Met Ala Phe Gly Thr Ser Ser Asp Val Glu Phe Asn Cys Thr Ala Asp Asn Leu Ala Ile Glu Gly Gly Met Ser Asp Leu Pro Ala Tyr Lys Leu Met Cys Phe Asp Ile Glu Cys Lys Ala Gly Gly Glu Asp Glu Leu Ala Phe Pro Val Ala Gly His Pro Glu Asp Leu Val Ile Gln Ile Ser Cys Leu Leu Tyr Asp Leu Ser Thr Thr Ala Leu Glu His Val Leu Leu Phe Ser Leu Gly Ser Cys Asp Leu Pro Glu Ser His Leu Asn Glu Leu Ala Ala Arg Gly Leu Pro Thr Pro Val Val Leu Glu Phe Asp Ser Glu Phe Glu Met Leu Leu Ala Phe Met Thr Leu Val Lys Gln Tyr Gly Pro Glu Phe Val Thr Gly Tyr Asn Ile Ile Asn Phe Asp Trp Pro Phe Leu Leu Ala Lys Leu Thr Asp Ile Tyr Lys Val Pro Leu Asp Gly Tyr Gly Arg Met Asn Gly Arg Gly Val Phe Arg Val Trp Asp Ile Gly Gln Ser His Phe Gln Lys Arg Ser Lys Ile Lys Val Asn Gly Met Val Asn Ile Asp Met Tyr Gly Ile Ile Thr Asp Lys Ile Lys Leu Ser Ser Tyr Lys Leu Asn Ala Val Ala

530 535 540

Glu Ala Val Leu Lys Asp Lys Lys Lys Asp Leu Ser Tyr Arg Asp Ile 545 550 550 560

- Pro Thr Tyr Tyr Ala Ala Gly Pro Ala Gln Arg Gly Val Ile Gly Glu
 565 570 575
- Tyr Cys Ile Gln Asp Ser Leu Leu Val Gly Gln Leu Phe Phe Lys Phe 580 585 590
- Leu Pro His Leu Glu Leu Ser Ala Val Ala Arg Leu Ala Gly Ile Asn 595 600 605
- Ile Thr Arg Thr Ile Tyr Asp Gly Gln Gln Ile Arg Val Phe Thr Cys 610 620
- Leu Leu Arg Leu Ala Asp Gln Lys Gly Phe Ile Leu Pro Asp Thr Gln 625 630 635 640
- Gly Arg Phe Arg Gly Ala Gly Gly Glu Ala Pro Lys Arg Pro Ala Ala 645 650 655
- Ala Arg Glu Asp Glu Glu Arg Pro Glu Glu Glu Glu Glu Asp Glu Asn 660 670
- Glu Arg Glu Glu Gly Gly Glu Arg Glu Pro Glu Gly Ala Arg Glu 675 680 685
- Thr Ala Gly Arg His Val Gly Tyr Gln Gly Ala Arg Val Leu Asp Pro 690 695 700
- Thr Ser Gly Phe His Val Asn Pro Val Val Phe Asp Phe Ala Ser 715 720
- Leu Tyr Pro Ser Ile Ile Gln Ala His Asn Leu Cys Phe Ser Thr Leu 725 730 735
- Ser Leu Arg Ala Asp Ala Val Ala His Leu Glu Ala Gly Lys Asp Tyr 740 745 750
- Leu Glu Ile Glu Val Gly Gly Arg Arg Leu Phe Phe Val Lys Ala His
 755 760 765
- Val Arg Glu Ser Leu Leu Ser Ile Leu Leu Arg Asp Trp Leu Ala Met 770 775 780
- Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala 785 790 795 800
- Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val Cys Asn Ser 805 810 815
- Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu Leu Pro Cys Leu His 820 825 830
- Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu Leu Ala Thr 835 840 845
- Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala 850 855 860

Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met 865 870 875 880

- Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly 885 890 895
- Leu Thr Ala Ala Gly Leu Thr Ala Val Gly Asp Lys Met Ala Ser His
 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Pro Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys 945 950 955 960
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr 1040 1045 1050
- Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala 1085 1090 1095
- Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala Leu Pro Ser 1100 1105 1110
- Pro Ala Lys Arg Pro Arg Glu Thr Pro Ser Pro Ala Asp Pro Pro 1115 1120 1125
- Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu Val Ser Glu Leu Ala 1130 1135 1140
- Glu Asp Pro Ala Tyr Ala Ile Ala His Gly Val Ala Leu Asn Thr 1145 1150 1155
- Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala Ala Cys Val Thr Phe 1160 1165 1170
- Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile Thr Glu Ser Leu Leu 1175 1180 1185

Lys Arg Phe Ile Pro Glu Val Trp His Pro Pro Asp Asp Val Ala 1190 1195 1200

Ala Arg Leu Arg Thr Ala Gly Phe Gly Ala Val Gly Ala Gly Ala 1205 1210 1215

Thr Ala Glu Glu Thr Arg Arg Met Leu His Arg Ala Phe Asp Thr 1220 1225 1230

Leu Ala 1235

<210> 19

<211> 1235

<212> PRT

<213> herpes simplex

<400> 19

Met Phe Ser Gly Gly Gly Pro Leu Ser Pro Gly Gly Lys Ser Ala 1 5 10 15

Ala Arg Ala Ser Gly Phe Phe Ala Pro Ala Gly Pro Arg Gly Ala 20 25 30

Gly Arg Gly Pro Pro Pro Cys Leu Arg Gln Asn Phe Tyr Asn Pro Tyr 35 40 45

Leu Ala Pro Val Gly Thr Gln Gln Lys Pro Thr Gly Pro Thr Gln Arg
50 55 60

His Thr Tyr Tyr Ser Glu Cys Asp Glu Phe Arg Phe Ile Ala Pro Arg 65 70 75 80

Val Leu Asp Glu Asp Ala Pro Pro Glu Lys Arg Ala Gly Val His Asp 85 90 95

Gly His Leu Lys Arg Ala Pro Lys Val Tyr Cys Gly Gly Asp Glu Arg 100 105 110

Asp Val Leu Arg Val Gly Ser Gly Gly Phe Trp Pro Arg Arg Ser Arg 115 120 . 125

Leu Trp Gly Gly Val Asp His Ala Pro Ala Gly Phe Asn Pro Thr Val
130 135 140

Thr Val Phe His Val Tyr Asp Ile Leu Glu Asn Val Glu His Ala Tyr 145 150 155 160

Gly Met Arg Ala Ala Gln Phe His Ala Arg Phe Met Asp Ala Ile Thr 165 170 175

Pro Thr Gly Thr Val Ile Thr Leu Leu Gly Leu Thr Pro Glu Gly His
180 185 190

Arg Val Ala Val His Val Tyr Gly Thr Arg Gln Tyr Phe Tyr Met Asn 195 200 205

Lys Glu Glu Val Asp Arg His Leu Gln Cys Arg Ala Pro Arg Asp Leu 210 220

Cys Glu Arg Met Ala Ala Ala Leu Arg Glu Ser Pro Gly Ala Ser Phe Arg Gly Ile Ser Ala Asp His Phe Glu Ala Glu Val Val Glu Arg Thr Asp Val Tyr Tyr Tyr Glu Thr Arg Pro Ala Leu Phe Tyr Arg Val Tyr Val Arg Ser Gly Arg Val Leu Ser Tyr Leu Cys Asp Asn Phe Cys Pro Ala Ile Lys Lys Tyr Glu Gly Gly Val Asp Ala Thr Thr Arg Phe Ile Leu Asp Asn Pro Gly Phe Val Thr Phe Gly Trp Tyr Arg Leu Lys Pro Gly Arg Asn Asn Thr Leu Ala Gln Pro Arg Ala Pro Met Ala Phe Gly Thr Ser Ser Asp Val Glu Phe Asn Cys Thr Ala Asp Asn Leu Ala Ile Glu Gly Gly Met Ser Asp Leu Pro Ala Tyr Lys Leu Met Cys Phe Asp Ile Glu Cys Lys Ala Gly Gly Glu Asp Glu Leu Ala Phe Pro Val Ala Gly His Pro Glu Asp Leu Val Ile Gln Ile Ser Cys Leu Leu Tyr Asp Leu Ser Thr Thr Ala Leu Glu His Val Leu Leu Phe Ser Leu Gly Ser Cys Asp Leu Pro Glu Ser His Leu Asn Glu Leu Ala Ala Arg Gly Leu Pro Thr Pro Val Val Leu Glu Phe Asp Ser Glu Phe Glu Met Leu Leu Ala Phe Met Thr Leu Val Lys Gln Tyr Gly Pro Glu Phe Val Thr Gly Tyr Asn Ile Ile Asn Phe Asp Trp Pro Phe Leu Leu Ala Lys Leu Thr Asp Ile Tyr Lys Val Pro Leu Asp Gly Tyr Gly Arg Met Asn Gly Arg Gly Val Phe Arg Val Trp Asp Ile Gly Gln Ser His Phe Gln Lys Arg Ser Lys Ile Lys Val Asn Gly Met Val Asn Ile Asp Met Tyr Gly Ile Ile Thr Asp Lys Ile Lys Leu Ser Ser Tyr Lys Leu Asn Ala Val Ala Glu Ala Val Leu Lys Asp Lys Lys Lys Asp Leu Ser Tyr Arg Asp Ile

Pro Ala Tyr Tyr Ala Ala Gly Pro Ala Gln Arg Gly Val Ile Gly Glu Tyr Cys Ile Gln Asp Ser Leu Leu Val Gly Gln Leu Phe Phe Lys Phe Leu Pro His Leu Glu Leu Ser Ala Val Ala Arg Leu Ala Gly Ile Asn Ile Thr Arg Thr Ile Tyr Asp Gly Gln Gln Ile Arg Val Phe Thr Cys Leu Leu Arg Leu Ala Asp Gln Lys Gly Phe Ile Leu Pro Asp Thr Gln Gly Arg Phe Arg Gly Gly Gly Glu Ala Pro Lys Arg Pro Ala Ala Ala Arg Glu Asp Glu Glu Arg Pro Glu Glu Glu Gly Glu Asp Glu Asp Glu Arg Glu Glu Gly Gly Glu Arg Glu Pro Glu Gly Ala Arg Glu Thr Ala Gly Arg His Val Gly Tyr Gln Gly Ala Arg Val Leu Asp Pro Thr Ser Gly Phe His Val Asn Pro Val Val Val Phe Asp Phe Ala Ser Leu Tyr Pro Ser Ile Ile Gln Ala His Asn Leu Cys Phe Ser Thr Leu Ser Leu Arg Ala Asp Ala Val Ala His Leu Glu Ala Gly Lys Asp Tyr Leu Glu Ile Glu Val Gly Gly Arg Arg Leu Phe Phe Val Lys Ala His Val Arg Glu Ser Leu Leu Ser Ile Leu Leu Arg Asp Trp Leu Ala Met Arg Lys Gln Ile Arg Ser Arg Ile Pro Gln Ser Ser Pro Glu Glu Ala Val Leu Leu Asp Lys Gln Gln Ala Ala Ile Lys Val Val Cys Asn Ser Val Tyr Gly Phe Thr Gly Val Gln His Gly Leu Leu Pro Cys Leu His Val Ala Ala Thr Val Thr Thr Ile Gly Arg Glu Met Leu Leu Ala Thr 8.45 Arg Glu Tyr Val His Ala Arg Trp Ala Ala Phe Glu Gln Leu Leu Ala Asp Phe Pro Glu Ala Ala Asp Met Arg Ala Pro Gly Pro Tyr Ser Met Arg Ile Ile Tyr Gly Asp Thr Asp Ser Ile Phe Val Leu Cys Arg Gly

885 890 895

- Leu Thr Ala Ala Gly Leu Thr Ala Val Gly Asp Lys Met Ala Ser His
 900 905 910
- Ile Ser Arg Ala Leu Phe Leu Ser Pro Ile Lys Leu Glu Cys Glu Lys 915 920 925
- Thr Phe Thr Lys Leu Leu Leu Ile Ala Lys Lys Lys Tyr Ile Gly Val 930 935 940
- Ile Tyr Gly Gly Lys Met Leu Ile Lys Gly Val Asp Leu Val Arg Lys 945 950 955 960
- Asn Asn Cys Ala Phe Ile Asn Arg Thr Ser Arg Ala Leu Val Asp Leu 965 970 975
- Leu Phe Tyr Asp Asp Thr Val Ser Gly Ala Ala Ala Ala Leu Ala Glu 980 985 990
- Arg Pro Ala Glu Glu Trp Leu Ala Arg Pro Leu Pro Glu Gly Leu Gln 995 1000 1005
- Ala Phe Gly Ala Val Leu Val Asp Ala His Arg Arg Ile Thr Asp 1010 1015 1020
- Pro Glu Arg Asp Ile Gln Asp Phe Val Leu Thr Ala Glu Leu Ser 1025 1030 1035
- Arg His Pro Arg Ala Tyr Thr Asn Lys Arg Leu Ala His Leu Thr 1040 . 1045 1050
- Val Tyr Tyr Lys Leu Met Ala Arg Arg Ala Gln Val Pro Ser Ile 1055 1060 1065
- Lys Asp Arg Ile Pro Tyr Val Ile Val Ala Gln Thr Arg Glu Val 1070 1075 1080
- Glu Glu Thr Val Ala Arg Leu Ala Ala Leu Arg Glu Leu Asp Ala 1085 1090 1095
- Ala Ala Pro Gly Asp Glu Pro Ala Pro Pro Ala Ala Leu Pro Ser 1100 1105 1110
- Pro Ala Lys Arg Pro Arg Glu Thr Pro Leu His Ala Asp Pro Pro 1115 1120 1125
- Gly Gly Ala Ser Lys Pro Arg Lys Leu Leu Val Ser Glu Leu Ala 1130 1135 1140
- Glu Asp Pro Ala Tyr Ala Ile Ala His Gly Val Ala Leu Asn Thr 1145 1150 1155
- Asp Tyr Tyr Phe Ser His Leu Leu Gly Ala Ala Cys Val Thr Phe 1160 1165 1170
- Lys Ala Leu Phe Gly Asn Asn Ala Lys Ile Thr Glu Ser Leu Leu 1175 1180 1185
- Lys Arg Phe Ile Pro Glu Val Trp His Pro Pro Asp Asp Val Ala 1190 1195 1200

Ala Arg Leu Arg Ala Ala Gly Phe Gly Ala Val Gly Ala Gly Ala 1205 1210 1215

Thr Ala Glu Glu Thr Arg Arg Met Leu His Arg Ala Phe Asp Thr 1220 1230

Leu Ala 1235